

EPA High Production Volume (HPV) Track Physical-Chemical End Point: Molting Point Spensor ID 110002: Albemarle Corporation Create Date 4.6/01CAS Number 3194556 Cyclododecane, 1,2.5,6,9,10-hexabromo-Study Number Consortia ID 1101012 CMA Brominated Flame Retardant Industry Panel (BFRIP) Completed: **Revision Date:** 12/5/01 **Test Substance** Remarks The test substance consisted of various commercial products. **Chemical Category Method** >> Method/Guideline followed Not specified. >> GLP Unknown >> Year study performed 1994 Remarks for Method Results >> Precision range >> Melting Point Value 175

195

>> Unit °C

>> Upper Value

EPA High Production Volume (HPV) Track Physical-Chemical End Point: Melting Point

| | | San | | |
|--------------------|---------------|---|----------------|---------|
| Sponsor ID | 1100021 | Albemarie Corporation | Greate Date | 1,6/01 |
| CAS Number | 3194550 | Cyclododecane, 1,2,5.6,9,10-hexabromo- | Study Number | : |
| Consortia ID | 1101012 | CMA Browinated Flame Retardant Industry Panel (BFRIP) | Completed: | Υ |
| >> Decomposition | on Yes | | | |
| >> Sublimation | No | | | |
| Results Remai | 1 1 | ing points have been reported for different products: 175 187-195 degrees C (Saytex-HM), 190 degrees C (GLCC | | (Saytex |
| Conclusions | HBCD is a so | lid at room temperature whose melting point varies with | composition. | |
| Data Quality | Reliability | Good | | |
| Data Reliability R | emarks | | | |
| | The melting p | oint data was provided by commercial manufacturers of | the substance. | |
| Reference | | | | |
| >> Remarks | IUCLID Datas | et. Substance ID: 25637-99-4. 18-Feb-2000. | | |
| | | | | |

EPA High Production Volume (HPV) Track Physical-Chemical End Point: Melting Point

| Sponsor ID | 1109021 | Albemarle Corporation | Create Date 1/6/01 |
|--------------|---------|---|--------------------|
| CAS Number | 3194556 | Cyclododecane, 1,2,5,6,9,10-hexabromo- | Study Number |
| Consortia ID | 1101012 | CMA Brominated Flame Retardant Industry Panel (BFRIP) | Completed: Y |
| General | | | |
| | | | |
| | | | |
| | | | |
| | | | |

| Sponser ID | 1100021 | Albemarle Corporation | Create Date |
|-----------------------------------|--|--|---|
| CAS Number | 3194556 | Cyclododecane. 1,2,5,6,9.10-hexabromo- | Study Number |
| Consortia ID | 1101012 | CMA Brominated Flame Retardant Industry Panel (BFRIP) | Completed: Y |
| | | | Revision Dat |
| t Substand | 20 | | 12/5/ |
| Remai | commercial pro Lakes Chemical homogeneity. T | was a composite of equal parts of the commercial heduct produced by Albemarle Corporation, Dead Seal Corporation. The test article composite was analyzing results of the analysis indicated the test article was apponents: HBCD beta isomer 8.5%, HBCD alpha isomer 8.5%, HB | Bromine Group, and Greated for characterization and as homogeneous and con |
| emical Categ | | | |
| hod Method/Guid OPPTS 830.7 | deline followed | ient (n-Octanol/Water), Generator Column Method | |
| hod Method/Guid OPPTS 830.7 | deline followed | ient (n-Octanol/Water), Generator Column Method >> Year study p | performed 1997 |
| hod Method/Guid | deline followed 7560 Partition Coeffici | >> Year study p | |
| hod Method/Guid | Remarks for Me A single generate Chromosorb W H substance in octation with octanol was Samples of the e | >> Year study p | column was packed with lution of the test nol were analyzed. The eagent water saturated elute the test substance. |

>> Value of Log Pow

>> Precision

5.625

EPA High Production Volume (HPV) Track Partition Coefficient

Physical-Chemical End Point:

| Sponsor ID | 1100021 | Albemarle Corporation | Create Date | 1 6/01 |
|-------------------------------|------------|---|--------------|--------|
| CAS Number | 3194550 | Cyclododecane, 1,2.5.6,9.10-hexabromo- | Study Number | 1 |
| Consortia ID | 1101012 | CMA Brominated Flame Retardant Industry Panel (BFRIP) | Completed. | Υ |
| | | | | |
| | | | | |
| >> Upper Value | | 0 | | |
| >> Upper Value >> Temperature | 25 degrees | 0 | | |

Results Remark

HBCD's water solubility was previously determined to be 0.0034 mg/L (Stenzel and Markley, 1997).

No interferences were observed at or above the limit of quantitation in the matrix blank sample. The percent recovery of the 1.00 and 10.0 ug HBCD/L matrix fortifications were 104 and 85%. The mean recovery was calculated at 95% of nominal.

The nominal flow rate of reagent water through the generator column was measured prior to the start of sample collection. Flow rates were also calculated based on the volume and collection time of each sample that was analyzed. The pump setting was 1.0 mL/min and the flow rate was measured at 1.0 mL/min. The calculated flow rates for samples averaged 0.87 mL/min and ranged from 0.86 to 0.87 mL/min.

The mean concentration of HBCD measured in the aqueous samples eluted from the generator column was 3.97 ug HBCD/L or 6.19 x 10-9 M (molecular weight of HBCD is 641.7 g/mole).

The mean concentration of HBCD measured in the octanol stock solution samples was 1.67 a HBCD/L or 2.61 x 10-3 M (molecular weight of HBCD is 641.7 g/mole).

Conclusions

The octanol/water partition (Kow) coefficient was calculated from the following equation:

Kow = Measured Concentration in Octanol (M)

Measured Concentration in Aqueous Samples (M)

Based on the results from octanol samples collected from the stock solution and aqueous samples collected from the generator column, the mean octanol/water partition coefficient (Kow) for HBCD was determined to be $4.22 \times 10-5$ (log Kow = 5.625).

Reliability High

Data Reliability Remarks

EPA High Production Volume (HPV) Track Partition Coefficient

Physical-Chemical End Point:

| Sponsor ID | 1100021 | Albemarle Corporation | Create Date | 1/6/01 |
|--------------|----------|---|--------------|--------|
| CAS Number | 319.3556 | Cyclododecane, 1,2.5,6,9.10-hexabromo- | Study Number | 1 |
| Consortia ID | 1101012 | CMA Brominated Flame Retardant Industry Panel (BFRIP) | Completed: | Y |

This study was performed according to current guidelines and Good Laboratory Practices by a laboratory with considerable experience with these studies. Extensive attention was paid to analytical method development and performance.

Reference

>> Remarks

MacGregor, J and Nixon, W. (1997) Hexabromocyclododecane (HBCD): Determination of n-Octanol/Water Partition Coefficient. Project Number: 439C-104. Wildlife International LTD. Easton, MD.

General

Study sponsored by the Chemical Manufacturers Association Brominated Flame Retardant Industry Panel, Arlington, VA.

| PA High | Production | | and the same of th | | | |
|----------------------|--|--|--|--|--|--|
| Sponsor ID | 1100021 | Albemaríe Corporation | | Create | Date | To the production of the contract of the contr |
| CAS Number | 3194556 | Cyclododecane, 1,2,5,6,9.1 |)-hexabromo- | Study | Number | |
| Con sortia ID | 1101012 | CMA Brominated Flame Re | tardant Industry Panel (BF | RIP) Compi | eted: | Y |
| | | | | | Revisio | n Date: |
| t Substanc | <u>Ce</u> | | | | | 12/5/01 |
| Remar | commercial prod Lakes Chemical homogeneity. T | was a composite of equal duct produced by Albeman Corporation. The test and the results of the analysis mponents: HBCD beta iso | te Corporation, Dead Sicle composite was and indicated the test articles. | Sea Bromine G alyzed for char e was homoge | Broup, an racteriza eneous a | nd Great tion and and contai |
| mical Catego | ory | | | | | |
| hod | _ | | | | | |
| | | | | | | |
| Method/Guid | deline followed | | | | | |
| | | TS 830 7950 Vapor Press | III | | | Management of the state of the |
| | | TS 830.7950 Vapor Press | urė | | · · · · · · · · · · · · · · · · · · · | |
| OECD Method | | TS 830.7950 Vapor Press | ure >> Year study | performed | 1997 | |
| DECD Method | | TS 830.7950 Vapor Press | | performed | 1997 | ************************************** |
| DECD Method | | | | performed | 1997 | |
| DECD Method | Remarks for Meta The objective of the temperature using | | >> Year study the vapor pressure of SRG). The SRG metho | HBCD at amb | pient | he |
| DECD Method | Remarks for Meta The objective of the temperature using extremely low vapon The SRG system of the statement | hod nis study was to determine a spinning rotor gauge (S | >> Year study the the vapor pressure of SRG). The SRG methor this substance. Inport 10 mL beaker in the | HBCD at amb od was chosen ne sample cha | pient due to t | 100000000000000000000000000000000000000 |
| DECD Method | Remarks for Meta The objective of the temperature using extremely low vapor and the SRG system was open to the value of the temperature of the tem | hod his study was to determine a spinning rotor gauge (Sor pressure anticipated for was configured with an er ents. The system baseling st substance in a 10 mL be used to monitor the stead accum pumps, and the propumps. The steady-stae | >> Year study the vapor pressure of SRG). The SRG methor this substance. Inpty 10 mL beaker in the pressure and out-gase eaker was placed in the y-state pressure of the essure increase from the | HBCD at amb od was chosen ne sample cha ssing rate were e sample char sample while ne sample while | oient due to to mber to e each mber. The the systele the val | make ne em ve |
| OECD Method GLP Yes | Remarks for Metal The objective of the temperature using extremely low vapar control measurement measured twice. A sample of the temperature was open to the values of the period of th | hod his study was to determine a spinning rotor gauge (Sor pressure anticipated for was configured with an er ents. The system baseling st substance in a 10 mL be used to monitor the stead accum pumps, and the propumps. The steady-stae | >> Year study the vapor pressure of SRG). The SRG methor this substance. Inpty 10 mL beaker in the pressure and out-gase eaker was placed in the y-state pressure of the essure increase from the | HBCD at amb od was chosen ne sample cha ssing rate were e sample char sample while ne sample while | oient due to to mber to e each mber. The the systele the val | make ne em ve |
| OECD Method GLP Yes | Remarks for Meta The objective of the temperature using extremely low vapor and the SRG system was used to the system was open to the value was closed to the system was repeated three systems. | hod his study was to determine a spinning rotor gauge (Sor pressure anticipated for was configured with an er ents. The system baseling st substance in a 10 mL be used to monitor the stead accum pumps, and the propumps. The steady-stae | >> Year study the vapor pressure of SRG). The SRG methor this substance. Inpty 10 mL beaker in the pressure and out-gase eaker was placed in the y-state pressure of the essure increase from the | HBCD at amb od was chosen ne sample cha ssing rate were e sample char sample while ne sample while | oient due to to mber to e each mber. The the systele the val | make ne em ve |
| | Remarks for Meta The objective of the temperature using extremely low vapor and the SRG system was used to the system was open to the value was closed to the system was repeated three systems. | hod his study was to determine a spinning rotor gauge (Sor pressure anticipated for was configured with an er ents. The system baseling st substance in a 10 mL be used to monitor the stead accum pumps, and the propumps. The steady-stae | >> Year study the vapor pressure of SRG). The SRG methor this substance. Inpty 10 mL beaker in the pressure and out-gase eaker was placed in the y-state pressure of the essure increase from the | HBCD at amb od was chosen ne sample cha ssing rate were e sample char sample while ne sample while | oient due to to mber to e each mber. The the systele the val | make ne em ve |

EPA High Production Volume (HPV) Track Vapor Pressure

Physical-Chemical End Point:

| Sponsor ID | [10002] | Albemarle Corporation | Create Date | 1-6/01 |
|--------------|---------|---|--------------|--------|
| CAS Number | 319 556 | Cyclododecane, 1,2.5,6,9,10-hexabromo- | Study Number | i |
| Consortia ID | 1101012 | CMA Brominated Flame Retardant Industry Panel (BFRIP) | Completed: | Υ |

| >> Upper Value | 0 |
|-------------------------|-------|
| >> Unit Pascals | |
| >> Temperature 21 degre | ees C |
| >> Decomposition No | |

Results Remark The baseline pressure of the system containing an empty beaker was determined to be less than 1 x 10E-7 Pa for both measurements. The technical specifications of the SRG indicated the low end of the measurement range to be 1 x 10E-5 Pa. The baseline pressure was considered to be essentially zero, and indicated the system was free of contamination. The outgassing rate (slope) was <1 x 10E-7 Pa/sec for both measurements. The out-gassing rate indicated there were no leaks in the system.

> The mean steady-state pressue of the HBCD sample was 6.166 x 10E-5 based on three separate determinations. The slope of the line fit to the pressue increase data was less than the out-gassing rate of the empty system for each determination, indicating the system had achieved saturation of the gas from the HBCD sample and was leak-free. The intercept was only slightly greater than the steady-state pressure. The temperature of the system averaged 21 degrees C.

The vapor pressure for each determination of the HBCD sample was calculated from the following equation:

Vapor Pressure = intercept of sample - mean intercept of empty system.

The mean vapor pressure of HBCD was determined to be 6.27 x 10E-5 Pa with a standard deviation of $0.21 \times 10E-5$.

The vapor pressures of di(2-ethyl-hexyl)phthalate and hexachlorobenzene were measured using the same SRG system and determined to be 4.3 x 10E-5 Pa and 1.6 x 10E-3 Pa, respectively. Both of these measurements were consistent with ranges found in the literature.

Conclusions

Physical-Chemical End Point: Vapor Pressure

| Cg | | VOIGITIES (111 V) 11 CCT Vapor Pressure | 8 |
|--------------------|------------------------------------|---|---|
| Sponsor ID | 1100021 | Albemarle Corporation | Create Date |
| CAS Number | 3194856 | Cyclododecane, 1,2,5,6,9,10-hexabromo- | Study Number |
| Consortia ID | 1101012 | CMA Brominated Flame Retardant Industry Panel (BFR) | P) Completed: Y |
| | Based on the re the vapor press | sults from three sets of measurements collected from the sets of measurements collected from the sets of HBCD was determined to be 6.27 x 10E-5 P | om the spinning rotor gauge a at 21 degrees C. |
| ata Quality | Reliability Hig | lh | |
| Data Reliability R | emarks | | |
| Reference | laboratory with o | performed according to current guidelines and Goo considerable experience with these studies. Extens d development and performance. | sive attention was paid to |
| >> Remarks | | ixon, W. (1997) Hexabromocyclododecane (HBC Using a Spinning Rotor Guage. Project Number: 4), Easton, MD. | |
| ieneral | | | |
| | Study sponsored Industry Panel. | by the Chemical Manufacturers Association Brom | inated Flame Retardant |
| | | | |

| Sponsor ID | | | Water Solubility | | |
|------------------|---|--|--|--|-----------------------------------|
| | [100021] | Albemarle Corporation | | Create Date | 1 (4/1) |
| CAS Number | 3194556 | Cyclododecane, 1,2,5,6,9,10-hexabroi | mo- | Study Number | |
| Consortia ID | 1101012 | GMA Brominated Flame Retardant Ind | dustry Panel (BFRIP) | Completed- | N |
| | | | | Revis | ion Date: |
| st Substance | | | | | 12/5/01 |
| Remarks | commercial pro Chemical Corpo homogeneity. 1 | was a composite of equal parts of the duct produced by Albemarle Corporation. The test article composite The results of the analysis indicated mponents: HBCD beta isomer 8.5% | oration, Dead Sea Browns was analyzed for chart the test article was | omine Group, a aracterization a homogeneous | nd Great La nd and containe |
| emical Category | | | | | |
| thod | | | | | |
| > Method/Guideli | ne followed | | | | |
| OECD Method 10 | 05, U.S. EPA 40 C | FR Ch. 1 Section 796.1860 Water | Solubility- Generato | r Column Metho | od |
| > GLP Yes | | | >> Year study per | formed | 1997 |
| | | ethod Deformed according to OECD Methor O Water Solubility- Generator Colum | | A 40 CFR Ch. | 1 |
| | A generator colu C and reagent w substance. Sam concentration of | mn was prepared. The column tenater was pumped through it at appriples of the eluate were collected at the test substance. The flow rate eximately half the original flow rate | roximately 2 mL per on analyzed to deter of reagent water thro | minute to elute mine the satura ugh the column | the test tion was |

EPA High Production Volume (HPV) Track Water Solubility

Physical-Chemical End Point:

| Sponsor ID | 1100021 | Albemarle Corporation | Create Date : 6/01 |
|-----------------------|-------------|---|--------------------|
| CAS Number | 3194556 | Cyclododecane, 1,2,5,6,9,10-hexabromo- | Study Number |
| Consortia ID | 1101012 | CMA Brominated Flame Retardant Industry Panel (BFRIP) | Completed: |
| >> Unit mg/L | | | |
| >> Temperature 25 c | degrees C | | |
| >> Solubility Categor | y Insoluble | | |
| | | | |
| | | | |
| >> pH Value | 8 | >> pKa Value 0 | |
| Poeulte Pomark | | | |

No interferences were observed at or above the limit of detection (0.5 ug HBCD/L) in any of the matrix blank or reagent blank samples. The peak area response for the matrix blanks was always below the response of the lowest calibration standard. The mean recovery from 10 matrix samples fortified at 10 ug/L was 105% (standard deviation 2.0), and ranged from 103% to 108%. The mean recovery from 10 matrix samples fortified at 1 ug/L was 104% (standard deviation 5.2),a nd ranged from 100% to 110%. The 1 ug/L concentration was considered the limit of quantiation.

A brief description of the analytical method is as follows: samples were extracted using dichloromethane (DCM). The DCM was evaporated to dryness and 1.0 ml o facetonitrile/water (50:50) was added. The samples was analyzed using HPLC/UV.

The nominal flow rate of reagent water through the generator column was initially set at 1.0 mL/min. The initial flow rate was measured at 2.0 mL/min prior to the start of sample collection. Samples were collected at this flow rate until the solubility plateau was achieved. The calculated flow rates for samples collected at the initial flow rate average 1.96 mL/min (range 1.88-1.98 mL/min). After the solubility plateau was achieved, the flow rate was reduced to ~half the initial flow rate. The reduced flow rate was measured at 1.0 mL/min prior to resuming sample collection. The calculated flow rates averaged 0.92 mL/min (range 0.91-0.93)

All samples collected at a nominal flow rate of 2.0 mL/min were analyzed and the solubility limit was considerated to have been achieved when at least 5 consecutive samples gave similar results. The mean concentration in samples meeting this criteria was 0.0034 mg/L with a standard deviation of 0.23.

The results from analyses of samples eluted at a nominal flow rate of 1.0 mL/min found a mean

Physical-Chemical End Point: Water Solubility

| C. A. Ting. | · · · · · · · · · · · · · · · · · · · | Volume (111 4) 11 GCN Water Solubility | | |
|--------------------|---------------------------------------|--|---------------------|----------|
| Sponsor ID | [100021] | Albemarle Corporation | Create Date | 4.6/0 |
| CAS Number | 3194550 | Cyclododecane, 1,2,5,6,9,10-hexabromo- | Study Number | |
| Consortia ID | 11011112 | CMA Brominated Flame Retardant Industry Panel (BFRIP |) Completed: | N |
| | HBCD concentra | ation of 0.0033 mg/L with a standard deviation of 0.2 | 20. | |
| | | ater obtained from Wildlife International Ltd's well in 3 (range 8.2-8.4). | February/March 19 | 97 had |
| | | as no ionizable groups and therefore the pKa value of the "pka value" field becuase this was a mandatory | | /alue of |
| Conclusions | | | | |
| | The solubility of | HBCD in water was determined to be 0.0034 +/- 0.2 | mg/L at 25 degree | s C. |
| Data Quality | Reliability Hig | h : | | |
| Data Reliability R | emarks | | | |
| | laboratory with o | performed according to current guidelines and Good considerable experience with these studies. Extension development and performance. | | |
| <u>leference</u> | | | | |
| >> Remarks | | Markley, B. (1997) Hexabromocyclododecane (HB0 Project Number: 439C-105. Wildlife International I | | of the |
| eneral | | | | |
| | Study sponsored Industry Panel, A | d by the Chemical Manufacturer's Association Brom Arlington, VA. | inated Flame Retard | dant |

Environmental Fate and Pathway End Point: Biodegradation

| Sponsor ID | 1100021 | Albemarle Corporation | Create Date | 1/6/01 |
|--------------|-----------|--|--------------|--------|
| CAS Number | 319456 | Cyclododecane. 1,2,5,6,9,10-hexabromo- | Study Number | i |
| Consortia ID | [1101012] | CMA Brondinated Flame Retardant Industry Panel (BFRIP) | Completed: | Ÿ |

Revision Date:

Test Substance

12/5/01

Remarks

The test article was a composite of equal parts of the commercial hexabromocyclododecane (HBCD) commercial product produced by Albemarle Corporation, Dead Sea Bromine Group, and Great Lakes Chemical Corporation. The test article composite was analyzed for characterization and homogeneity. The results of the analysis indicated the test article was homogeneous and contained the following components: HBCD beta isomer 8.5%, HBCD alpha isomer 6.0%, HBCD gamma isomer 79.1%.

Chemical Category

Method

EPA OPPTS Method 835.3200: Ready Biodegradability, Closed Bottle Test; OECD Guideline 301D

>> Test Type

aerobic

>> GLP Yes

>> Year study performed

1996

>> Contact Time

28

>> inoculum

activated sludge, domestic, adapted

Remarks for Method

The test contained an inoculum control group, a reference group and a treatment group. The blank control, reference, and treatment groups contained ten replicate test chambers. The inoculum control was used to measure the dissolved oxygen consumption of the inoculum and was not dosed with a carbon source. The reference chambers were dosed with sodium benzoate, a substance known to be biodegradable, at a concentration of 2 mg/L. The treament group test chambers were used to evaluate the test substance at 7.7 mg/L. Measurements of oxygen consumption were performed on two test chambers from the control, reference and treatment groups on days 0, 7, 14, 21, and 28.

The test inoculum was secondary clarifier supernatant collected from Prospect Bay Wastewater Treatment Facility, Grasonville, MD. The theoretical oxygen demand value used to calculate the percent degradation of the test substance was 0.75 mg O2/mg.

<u>Results</u>

Environmental Fate and Pathway End Point:

| EPA High Pr | oduction | VOIUME (MPV) I Tack Blodegradation | |
|---------------------|--|--|--------------------------|
| Sponsor ID | 1100024 | Albemarle Corporation | Create Date (i, o/0) |
| CAS Number | 3194,556 | Cyclododecane: 1,2 5,6.9 10-hexabromo- | Study Number 1 |
| Consortia ID | 1101012 | CMA Brominated Flame Retardant Industry Panel (BFRIP) | Completed: Y |
| >> Precision = | | | |
| >> Degradation Valu | 10 | 0 | |
| >> Upper value | | 0 | |
| >> Time Frame | | 28 | |
| >> Time Units Days | | | |
| >> Breakdown prod | ucts No | | |
| Results Remarks | g (COLUMN TO THE | | |
| | | e range recorded during the test was 18-20 degrees ount performed on the inoculum was $3.7 \times 10E4$ CFU | |
| | measured at 0, | /gen uptake exhibited by the control, reference, and 7, 14, 21 and 28 days. The oxygen depletion of the i 1.5 mg O2/L. Degradation of the test substance was | noculum control was less |

day test period.

The viability of the inoculum and validity of the test was supported by the results of the reference substance, sodium benzoate, degrading approximately 94%. An average percent biodegradation of > 60% was achieved by day 7, thereby fulfilling the criteria for a valid test.

Conclusions

Degradation of the test substance, HBCD, at 7.7 mg/L was not observed over the 28-day test period.

Environmental Fate and Pathway End Point: Biodegradation

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|--------------------|--|---|----------------------|-------|
| Spensor ID | 1160021 | Albemarie Corporation | Create Date | 4/69 |
| CAS Number | 3194556 | Cyclododecane 1,2,5,6,9.10-hexabromo- | Study Number | |
| Consortia ID | 1101012 | CMA Bronnnated Flame Retardant Industry Panel (BFRIP |) Completed: | Υ |
| Data Quality | Reliability Hig | gh | | |
| Data Reliability R | emarks | | | |
| | laboratory with | performed according to current guidelines and Good considerable experience with these studies. Extens od development and performance. | | |
| Reference | And the state of t | | | |
| >> Remarks | | d Haberlein, D. (1996) Hexabromocyclododecane (H 9E-102. Wildlife International Ltd. Easton, MD. | IBCD): Closed Bottle | Test. |
| | | | | |

Environmental Fate and Pathway End Point: Transport between Environmental Compartments (Fugacity)

| CAS Number Study Number Study Number | ra riigii | ı Fr | oduc Hon | volume (FIFV) Truck | between Environm | nental Compartments (Fugacity |
|--|--|--|--------------------|--------------------------------------|----------------------|-------------------------------|
| Complete Y Complete Y Complete Y Complete Y | Sponsor ID | | 3100021 | Albeniarle Corporation | | Create Date 4 (|
| Remarks Hexabromocyclododecane (HBCD) 12/18/01 | CAS Number | | 3181556 | Cyclododecane: 1,2,5,6,9,10-hexabror | no- | Study Number |
| Remarks Hexabromocyclododecane (HBCD) | Consortia ID | and the state of t | 1101012 | CMA Brominated Flame Retardant ind | fustry Panel (BFRIP) | Completed: |
| Remarks Hexabromocyclododecane (HBCD) mical Category thod Method/Guideline followed Developed by D. Mackay and co-workers Test Type Level III fugacity model Remarks for Method Model Used: Level III Fugacity Model (Full-Output), EPIWIN V3.04 Input parameters: chemical structure only; model default parameters accepted; model basd on emissions of 1000 kg/hr each to air, water and soil. Media 1. 0.000685%; Water: 2.06%; Soil: 40.1%; Sediment: 57.9% | | | | | | Revision Date: |
| Hexabromocyclododecane (HBCD) | st Substan | ce | | | | 12/18/01 |
| Method/Guideline followed Developed by D. Mackay and co-workers Test Type | | | Hexabromocycl | ododecane (HBCD) | | |
| Method/Guideline followed Developed by D. Mackay and co-workers Test Type Level III fugacity model >> Year study performed 2001 Remarks for Method Model Used: Level III Fugacity Model (Full-Output), EPIWIN V3.04 Input parameters: chemical structure only; model default parameters accepted; model basd on emissions of 1000 kg/hr each to air, water and soil. Media 1. 0.000685%; Water: 2.06%; Soil: 40.1%; Sediment: 57.9% Distribution Concentration | | | | | | |
| Method/Guideline followed Developed by D. Mackay and co-workers Test Type Level III fugacity model >> Year study performed 2001 Remarks for Method Model Used: Level III Fugacity Model (Full-Output), EPIWIN V3.04 Input parameters: chemical structure only; model default parameters accepted; model basd on emissions of 1000 kg/hr each to air, water and soil. Uits Media Distribution Concentration | | | | | | |
| Method/Guideline followed Developed by D. Mackay and co-workers Test Type | miss! Cst-s- |) | | | | |
| Developed by D. Mackay and co-workers Test Type | | ory | | | | |
| Developed by D. Mackay and co-workers Test Type | | delin | e followed | | | |
| Test Type | ······································ | | | n-workers · | | |
| Remarks for Method Model Used: Level III Fugacity Model (Full-Output), EPIWIN V3.04 Input parameters: chemical structure only; model default parameters accepted; model basd on emissions of 1000 kg/hr each to air, water and soil. Buits Media Distribution Concentration | Болоюро | , . | . maokay ana o | | | |
| Remarks for Method Model Used: Level III Fugacity Model (Full-Output), EPIWIN V3.04 Input parameters: chemical structure only; model default parameters accepted; model basd on emissions of 1000 kg/hr each to air, water and soil. Buits Media Distribution Concentration | Test Tyne | l ave | al III fugacity mo | dal | >> Vear et | idy performed 2001 |
| Model Used: Level III Fugacity Model (Full-Output), EPIWIN V3.04 Input parameters: chemical structure only; model default parameters accepted; model basd on emissions of 1000 kg/hr each to air, water and soil. Media 1. 0.000685%; Water: 2.06%; Soil: 40.1%; Sediment: 57.9% Distribution Concentration | , | | | | - 1 Gar Stt | lay periorinea 2001 |
| Input parameters: chemical structure only; model default parameters accepted; model basd on emissions of 1000 kg/hr each to air, water and soil. Bults Media C. 0.000685%; Water: 2.06%; Soil: 40.1%; Sediment: 57.9% Distribution Concentration | | _ | | | | |
| emissions of 1000 kg/hr each to air, water and soil. Bults Media C 0.000685%; Water: 2.06%; Soil: 40.1%; Sediment: 57.9% Distribution Concentration | | ٨ | flodel Used: Lev | el III Fugacity Model (Full-Output), | EPIWIN V3.04 | |
| Media The contraction Distribution Concentration | | | | | efault parameters ac | cepted; model basd on |
| Media : 0.000685%; Water: 2.06%; Soil: 40.1%; Sediment: 57.9% Distribution Concentration | | е | missions of 100 | o kg/nr each to air, water and soil. | | |
| Media : 0.000685%; Water: 2.06%; Soil: 40.1%; Sediment: 57.9% Distribution Concentration | | <u> </u> | | | | |
| : 0.000685%; Water: 2.06%; Soil: 40.1%; Sediment: 57.9% Distribution Concentration | | | | | | |
| Distribution Concentration | | ************* | | | | |
| | r: 0.000685%; | ; Wate | er: 2.06%; Soil: 4 | i0.1%; Sediment: 57.9% | | |
| Not provided by model. | Distribution | Conc | entration | | | |
| | | | Not provided t | ov model. | | |
| | | | o. p. oridod b | , , | | |
| | | | | | | |
| | | | | | | |

Environmental Fate and Pathway End Point: Transport between Environmental Compartments (Fugacity)

| | | VOIUME (HPV) I CACK between Environn | nental Compartments (F | ugacity / |
|--------------|--|--|---|-----------------------------------|
| Spensor ID | 11000021 | Albemarle Corporation | Create Date | 4-6/0 |
| CAS Number | 3194556 | Cyclododecane, 1,2,5,6,9.10-hexabromo- | Study Number | |
| Consortia ID | 1101012 | CMA Brominated Flame Retardant Industry Panel (BFRIP) | Completed: | |
| Results Rema | ırk | | | |
| nclusions | Liquid VP: 5 Melting Pt: 1 Log Kow: 7. | 25 x 10+7 sure: 1.68 x 10-8 mmHg .74 x 10-7 mm Hg (super-cooled) 80 deg C | | |
| | sediment (aptrace (appradvected. The model weither 0 or 10 predicted HE reacted. If reacted solutions to both total reacted 80% to soil a | t equal rates to air, water and soil, HBCD is predicted to opr. 58%) and soil (appr. 40%). Only appr. 2% would part of the proof of the | artition to water with of with only appr. 11% and soil emission rately to air, the model to sediment; 97% ont; total reacted = 710%. If released at element and one-third to BCD would partition to water and soil, HBC | only es as %. If qual soil; appr. |
| | | e above, HBCD is not expected to move from water, soil HBCD is not expected to move from soil into water. | or sediment to air. | |
| ata Quality | Reliability | High | | |

Reference

EPA High Production Volume (HPV) Track Environmental Fate and Pathway End Point: Transport between Environmental Compartments (Fugacity)

| Spensor ID | 1100021 | Albemarle Corporation | Create Date | 1,6/01 |
|----------------|--|---|-----------------------|--------|
| CAS Number | 3191956 | Cyclododecane, 1,2,5,6,9,10-hexabromo- | Study Number | i |
| Consortia ID | 1101017 | CMA Brominated Flame Retardant Industry Panel (BFRI | P) Completed: | Y |
| >> Remarks | Level III Fugad | city Model, EPIWIN V3.04, Syracuse Research Co | orporation, Syracuse, | NY. |
| | | | | |
| | | | | |
| <u>General</u> | The state of the s | | | |
| | | | | |
| | | | | |
| | | | | |

>> Analytical monitoring | HPLC/UV/VIS Detector; LOQ=0.04 ug/l

None needed - no mortality observed.

96 hours

Ecotoxicity End Point: Acute Toxicity to Fish

| | odderion | volume (iii v) irack | Acute Toxicity to Fish | | |
|------------------------|-----------------------------------|--|---|---------------------------------------|---|
| Sponsor ID | 1100021 | Albemarle Corporation | C | reate Date | -1.6.01 |
| CAS Number | 1104.56 | Cyclododecane, 1,2.5,6,9,10-hexabron | no- St | udy Numbe | r . |
| Consortia ID | 1101012 | CMA Brominated Flame Retardant Ind | ustry Panel (BFRIP) Co | ompleted: | N |
| · · | | | | | Revision Date |
| <u> Fest Substance</u> | | | | | 12/5/0 |
| Chemical Category | Great Lakes Che and homogeneit | cial product produced by Albemarle emical Corporation. The test article y. The results of the analysis indica lowing components: HBCD beta iso 9.1%. | composite was analyze ted the test article was | ed for char homogene | acterization eous and |
| >> Method/Guidelin | e followed | | | | |
| OECD Method 203 | | | | , , , , , , , , , , , , , , , , , , , | HILL THE STATE OF |
| >> Test Type | | | | | |
| flow-through | | | | | |
| >> GLP Yes | | | >> Year study perform | ned 1997 | 7 |
| >> Species | | | | | |
| Oncorhynchus my | rkiss | | | | |

Remarks for Method

>> Statistical Method

>> Exposure period

This study was performed according to OECD Method 203 and TSCA Title 40 of CFR, Part 797, Section 1400. Rainbow trout were exposed to one of five test concentrations, a solvent control, or the negative (well water) control. Two replicate test chambers were maintained in each treatment and control group. Ten rainbow trout were used in each test chamber for a total of 20 rainbow trout per test concentration. Nominal test concentrations were selected in consultation with the Sponsor, and were based on the solubility of the test compound in water (3.4 ug/L) and the results of an expoloratory rangefinding test. Due to co-eluting artifacts at 96

EPA High Production Volume (HPV) Track Acute Toxicity to Fish

Ecotoxicity End Point:

| Sponsor (D | 1400031 | Albemarle Corporation | Create Date | 4.6/01 |
|--------------|-----------|--|--------------|--------|
| CAS Number | 319 (550) | Cyclododecane. 1,2,5,6,9.10-hexabromo- | Study Number | 1 |
| Consortia ID | 110101? | CMA Brommated Flame Retardant Industry Panel (BFRIP) | Completed: | N |

hrs, mean measured test concentrations were determined analytically from samples of test water collected from each treatment and control group at the beginning of the test and at approximately 48 hrs.

The selection of exposure concentrations took into consideration the water solubility limit and a finding of no acute toxicity from an exploratory rangefinding test. The water solubility limit was determined in a generator column elution study to be 3.4 ug/L. However, there was a potential to have a slight enhancement of HBCD's water solubility due to the use of dimethylformamide (DMF) as a vehicle in the diluter system. For this reason, the highest test concentration selected for the acute toxicity test was twice the defined solubility limit (i.e., 6.8 ug/L). The series of 5 nominal test concentrations used in the test were 1.5, 2.2, 3.2, 4.6 and 6.8 ug/L. In this way, the solubility limit of HBCD was bracketed by the five concentrations.

Delivery of the test substance was initiated approximately 6 days prior to the introduciton of the fish to the test water in order to achieve equilibrium of the test substance in the test chambers. The fish wree indiscriminately assigned to exposure chambers at test initiation. Observations of mortality and other clinical signs were made approximately 1, 24, 48, 72 and 96 hrs after test initiation. The no mortality concentration and no observed effect concentration (NOEC) were determined by visual interpretation of the mortality and clinical observation data.

All fish were from the same source and year class, and the total length of the longest fish was no more than twice the length of the shortest. The average length of 10 negative control fish at the end of the test was 55 mm with a range of 50-61 mm. The wet weight of 10 negative control fish at the end of the test was 2.4 g with a range of 1.6-3.6 g. Loading, defined as the total wet weight of ifsh per liter of test water that passed through the test chamber in 24 hrs, was 0.27 g fish/L/day.

Temperature, dissolved oxygen, and pH were measured. Temperatues were within the limits of the 12 +/- 2 degrees C range established for the test. Dissolved oxygen concentrations were greater than or = 78% of saturation throughout the test. Water pH ranged from 8.2-8.3. Total organic carbon values were <1.0 mg C/L at test initiation and termination.

Test substance concentrations were determined via HPLC using a UV/VIS detector.

| F | 20 | 2 | 11 | ı | te |
|----------|----|---|----|---|----|
| <u> </u> | | v | w | | w |

| (esuits | | |
|---------------------------|--|--|
| >> Nominal concentration | 0, 0.0015, 0.0022, 0.0032, 0.0046, 0.0068 | |
| >> Measured concentration | 0, 0.00075, 0.0015, 0.0023, 0.0023, 0.0025 | |
| >> Precision > | | |
| >> Endpoint Type LC0 | | |

Ecotoxicity End Point: Acute Toxicity to Fish

| Sponsor ID (19902 | Albemarle Corporation | Create Date 46/01 |
|---------------------------------|--|-------------------|
| CAS Number 319.455 | . Cyclododecane: 1,2 5,6,9,10-hexabromo- | Study Number 1 |
| Consortia ID 110101. | CMA Brominated Flame Retardant Industry Panel (BFR | HP) Completed: N |
| >> Endpoint Value | 0 >> Unit used mg/L | |
| >> Concentration Type Nomina | >> Endpoint Time | 96 |
| >> Statistical results | | |
| None needed - no mortality obse | erved. | |
| Results Remark | | |

One set of pretest water samples was collected from the highest and lowest test concentrations and analyzed for HBCD concentrations. All pretest samples yielded concentrations that were considerably lower than the expected concentrations. The toxicity test was initiated and measurements of the HBCD concentrations in all test chambers were made at the beginning, middle and end of the test. In general, concentrations of HBCD made on samples collected at Day 0 and Day 2 were variable and failed to correspond to the dilution series expected from the nominal concentrations. All diluter operational records were checked and no evidence of any malfunctions or errors were found. Concentrations measured in the Day 4 samples were artificially high due to co-eluting artifacts at the retention time of HBCD. Attempts were made to separate the co-eluting artifacts during a reanalysis of the orginal Day 4 sample extracts, but the resulting chromatography showed those same interferences.

While the pattern of measured HBCD was unexpected, the results suggest that the exposure solutions were at the solubility limit of HBCD in the diluter system. The variability in the measured concentrations could have been influenced by the temperature of the exposure water (12 degrees C), the flow-through design, or the hydrophobic nature of HBCD (as evidenced by its nonpolar alkane structure and extremely low water solubility). These factors could explain both the failure of the measured values to correspond to the nominal concentrations and the variability observed in the measured concentrations. Overall, it appears that the solubility limit of HBCD, under the conditions that it was applied in this test, is within the range of 2.0 - 3.0 ug/L. The values obtained in the Day 4 samples were not reflective of the true conditions due to the co-eluting artifacts, and therefore, were not used in the study.

Temperatures were within the limits of the 12 + /-2 degrees C range established for the test. Dissolved oxygen concentration of > or = 78% of saturation were observed throughout the test. Water pH was consistent with values for moderately-hard water and ranged from 8.2 to 8.3. Total organic carbon values were < 1.0 mg C/L at test initiation and termination.

Observations for mortality and other signs of toxicity were made daily. Rainbow trout in the negative control and solvent control groups appeared healthy and normal throughout the test. All rainbow trout in the 1.5, 2.2, 3.2, 4.6 and 6.8 ug/L (nominal) treatment groups also appeared normal throughout the test with no mortalitites or overt signs of toxicity. Based on these results, the LC50 values at 24, 48, 72 and 96 hours were estimated to be >6.8 ug/L, the highest concentration tested.

Ecotoxicity End Point: Acute Toxicity to Fish

| Sponsor ID | 1199021 | Albemaric Corporation | Create Date | -3+6/0) |
|-----------------------|----------|---|--------------|---------|
| CAS Number | 31%15%t- | Cyclododecane, 1,2,5,6,9,10-hexabromp- | Study Number | 1 |
| Con sort ia ID | 1100012 | CMA Brominated Flame Retardant Industry Panel (BFRIP) | Completed | N |

Conclusions

The 96-hour LC50 value for rainbow trout exposed to HBCD was >6.8 ug/L (nominal) (>2.5 ug/L mean measured concentration), the highest concentration tested and twice HBCD's water solubility (3.4 ug/L). Based on the mortality and observation data, the 96-hour no morality concentration and the no-oberved-effect-concentration were 6.8 ug/L (nominal) (2.5 ug/L mean measured concentration) and was higher than the water solubility of HBCD.

| Data | Qua | lity |
|------|-----|------|
| | | |

Reliability

High

Data Reliability Remarks

This study was performed according to current guidelines and Good Laboratory Practices by a laboratory with considerable experience with these studies. Extensive attention was paid to analytical method development and performance.

Reference

>> Remarks

Graves, W and Swigert, J. (1997) Hexabromocyclododecane (HBCD): A 96-Hour Flow-Through Acute Toxicity Test with the Rainbow Trout (Oncorhynchus mykiss). Project Number: 439A-101. Wildlife International LTD, Easton, MD.

General

Study sponsored by the Chemical Manufacturers Association Brominated Flame Retardant Industry Panel, Arlington, VA.

Ecotoxicity End Point:
Acute Toxicity to Aquatic Invertebrates

| Sponsor ID | 1100021 | Albertarle Corporation | Create Date | 4/6/01 |
|--------------|---------|---|--------------|--------|
| CAS Number | 3101016 | Cyclododecane. 1,2,5,6,9,10-hexabromo- | Study Number | 1 |
| Consortia ID | 1101012 | CMA Brominated Flame Retardant Industry Panel (BFRIP) | Completed: | N |
| | | | | |

Revision Date:

Test Substance

12/5/01

Remarks

The test article was a composite of equal parts of the commercial hexabromocyclododecane (HBCD) commercial product produced by Albemarle Corporation, Dead Sea Bromine Group, and Great Lakes Chemical Corporation. The test article composite was analyzed for characterization and homogeneity. The results of the analysis indicated the test article was homogeneous and contained the following components: HBCD beta isomer 8.5%, HBCD alpha isomer 6.0%, HBCD gamma isomer 79.1%.

Chemical Category

| | 48. | |
|------|-----|--------|
| | ath | \sim |
| 11.1 | em | DO |
| | | |

| >> Met | hod/Guideline folk | owed | | |
|---------|--------------------|--|-------------------------|--|
| OEC | D Method 202; TSC | A Title 40 CFR, Part 797, Section 1300 | | 117 / 118 / 118 (118 / 118 / 118 / 118 / 118 / 118 / 118 / 118 / 118 / 118 / 118 / 118 / 118 / 118 / 118 / 118 |
| >> Tes | t Type | , | | |
| flow- | through | | | |
| >> GLF | Yes | | >> Year study performed | 1997 |
| >> Spe | cies nnia magna | | | |
| >> Ana | lytical monitoring | HPLC; Limit of Quantitation=0.4 ug/L | | |
| >> Exp | osure period | 48 Hours | | |
| >> Stat | istical Method | None - no dose response pattern | | |

Remarks for Method

Daphnids were exposed to one of five test concentrations, a solvent control or the negative (well water) control. Two replicate test chambers were maintained for each treatment and control group. Ten daphnids were used in each test chamber for a total of 20 daphnids per test concentration. Nominal test concentrations were based upon the solubility of the test substance in water (3.4 ug/L) and the results of an exploratory rangefinding toxicity test. Nominal test concentrations were 1.5, 2.2, 3.2, 4.6 and 6.8 ug/L. Mean measured test concentrations were analytically determined (HPLC with UV/VIS detector) from samples of test water collected from each treatment and control group at the beginning and end of the test.

Results

EPA High Production Volume (HPV) Track Acute Toxicity to Aquatic Invertebrates

Ecotoxicity End Point:

| Spansor ID | 1100021 | Albemarle Corporation | Create Date | 4-6/01 |
|--------------|----------|---|--------------|--------|
| CAS Number | 310 1556 | Cyclododecane, 1,2,5,6,9,10-hexabromo- | Study Number | ! |
| Consortia ID | 1101012 | CMA Brominated Flame Retardant Industry Panel (BFRIP) | Completed: | N |

Delivery of the test substance was initiated approximately 4 days prior to the introduction of the daphnids to the test water in order to acheive equilbrium of the test substance in the test chambers. Daphnids were indiscriminately assigned to exposure chambers at test initiation. Observations of mortality/immobility and other clinical signs were made approximately 2, 24 and 48 hours after test initiation. Cummulative percent mortality and immobility observed in the treatment groups were used to estimate EC50 values at 24 and 48 hours. The no mortality/immobility concentration and the no-observed-effect concentration (NOEC) were determined by visual interpretation of the mortality, immobility and clinical observation data.

Daphnid neonates used in the test were less than 24 hours old and were obtained from cultures maintained by Wildlife International Ltd, Easton, MD. Adult daphnids were cultured in water from the same source and at approximately the same tempreature as that used during the test except supplemented with selenium. Daphnids in the cultures were held for 15-29 days prior to collection of the juveniles for testing. The progency of 7 adults were used in the test. The adults were fed prior to test initiation, but neonates were not fed during the test. During the 14-day holding period preceeding the test, water temperatures ranged from 20.2 to 21.4 degrees C. The pH of the water ranged from 8.0 to 8.5. Disolved oxygen ranged from 8.2 to 9.0 mg/L.

A continuous-flow diluter was used to deliver each concentration of the test substance, a solvent control, and a negative (dilution water) control. Syringe pumps (Harvard Apparatus) were used to deliver the five test substance stock solutions and the solvent for the solvent control into mixing chambers assigned to each treatment level and the solvent control. The stock solutions were diluted with well water in the mixing chambers in order to obtain the desired test concentrations. The flow of dilution water to the mixing chambers was controlled by rotameters. Rotameters were calibrated prior to test initiaiton. The flow of test water from each mixing chamber was split and allowed to flow into replicate test chambers. The proportion of test water that was split into each replicate was checked prior to the test to ensure that flow rates varied by no more than +/- 10% of the mean for the two replicates.

The diluter was adjusted so that each test chamber received ~14 volume additions of test water every 24 hours. The stock solution delivery pumps were calibrated before the test, and the general operation of the diluter was checked visually at least two times daily during the test and once at the end of the test.

Test compartments were constructed from 300 mL glass beakers ~ 8 cm in diameter and 13 cm in height. The beakers were suspended in 8-L stainless steel test chambers filled with ~6.5 L of test water. Test chambers were indiscriminately positioned in a temperature-controlled water bath designed to maintain a temperature of 20+/-1 degreeC. The water bath was enclosed in a plexiglass ventilation hood. Test chambers were labeled with the project number, test concentration, and replicate.

The water used for culturing and testing was freshwater obtained from a well ~45 meters deep located on the Wildlife International Ltd. Site. The well water is characterized as moderatelyhard water. The dissolved oxygen content of the water ranged from 8.8-8.9, 9.0-9.1, and 8.8-8.9 mg/L at 0, 24, and 48 hours, respectively. The pH of the water was 8.1, 8.2-8.4, and 8.2-

Ecotoxicity End Point:
Acute Toxicity to Aquatic Invertebrates

| | | 40 L TL t | 1,6 40 | |
|--------------|----------|---|--------------|--------|
| Consortia ID | 1101012 | CMA Brominated Flame Retardant Industry Panel (BFRIP) | Completed: | N |
| CAS Number | 317 1556 | Cyclododecane. 1.2,5.6.9.10-hexabromo- | Study Number | 1 |
| Spensor ID | 1100021 | Albemaric Corporation | Create Date | 1/6/01 |

8.3 at 0, 24 and 48 hours, respectively. The temperature of the water ranged from 19.8-19.9 and 19.9-20.0 at 0 and 48 hours, respectively. The 0-hour dilution water measurements for hardness, alkalinity and specific conductance were 132 mg/L as CaC03, 176 mg/L as CaC03 and 320 umhos/cm, respectively.

Lighting was provided by fluorescent tubes that emitted wavelengths similar to natural sunlight. A photoperiod of 16 hours of light and 8 hours of darkness was controlled with an automatic timer. Light intensity at test initiation was ~ 242 lux at the surface of the water.

| >> Statistical results Statistics not performed due | to lack of dose repsonse. | | |
|---|------------------------------|---------------------|----|
| >> Concentration Type Nor | ninal >> Endpoint | Time | 48 |
| >> Endpoint Value | 0 | >> Unit used mg/L | |
| >> Endpoint Type LC0 | | | |
| >> Precision > | | | |
| >> Measured concentration | 0, 0.0024, 0.0018, 0.0021, 0 | 0.0023, 0.0032 mg/L | |
| | 0, 0.0010, 0.0022, 0.0002, 0 | | |
| >> Nominal concentration | 0, 0.0015, 0.0022, 0.0032, 0 | 0046, 0.0068 mg/L | |

Results Remark

The selection of exposure concentrations took into consideration the water solubility limit (3.4 ug/L) and a finding of no acute toxicity from an exploratory range finding test. However, there was a potential to have a slight enhancement of HBCD's water solubility due to the use of dimethyl formamide (DMF) as a vehicle in the diluter system. For this reason, the highest test concentration selected was twice the defined solubility limit (i.e., 6.8 ug/L). The series of nominal test concentrations bracketed the solubility limit of HBCD by five concentrations.

Two sets of pretest samples were collected from the highest and lowest test concentrations and analyzed. The Day -3 and -2 samples indicated that the test concentrations were stable, but somewhat lower than expected. Measurements of HBCD concentration in all test chambers were made at the beginning and end of the test. These measurements indicated that HBCD concentrations were generally similar across all treatment levels, and may reflect a

EPA High Production Volume (HPV) Track Acute Toxicity to Aquatic Invertebrates

Ecotoxicity End Point:

| Sponsor ID | 1160021 | Albemarle Corporation | Create Date | 4/6/01 |
|--------------|----------|---|--------------|--------|
| CAS Number | \$194456 | Cyclododecane 1,2,5.6,9,10-hexabromo- | Study Number | 1 |
| Consortia ID | 1101017 | CMA Brominated Flame Retardant industry Panel (BFRIP) | Completed: | N |

phenomenon in the delivery system whereby HBCD adsorbed to the physical surfaces of the diluter system. This could be due to the hydrophobic nature of HBCD as evidenced by its nonpolar alkane structure and extremely low water solubility. This characteristic could have enabled the inert surfaces (e.g. Stainless steel and Teflon) of the diluter system to eventually become saturated with HBCD. As this process progressed, an equilibium was established. The result of this new equilibrium was that concentrations of HBCD in the dilution water were approximately the solubility of HBCD in well water under flow-through conditions.

Dissolved oxygen concentrations of > or = 97% of saturation were observed throughout the test. Water pH ranged from 8.1-8.4. Total organic carbon in the dilution water at test initiation was <1.0 mg C/L.

Daily observations during the test showed that daphnids in the negative control and solvent control groups appeared healthy and normal. With the exception of one aberrant mortality in the 4.6 ug/L (nominal) treatment group, all daphnids in all treatment groups appeared normal throughout the test with no mortalities or overt signs of toxicity. Based on these results, EC50 values for 24 and 48 hours were estimated to be > 6.8 ug/L (nominal), the highest concentration tested.

Conclusions

HBCD was not acutely toxic to Daphnia magna. The 48-hour EC50 value for daphids exposed to HBCD was > 6.8 ug/L (nomimal) (>3.2 ug/L mean measured concentration), the highest concentration tested and twice HBCD's water solubility (3.4 ug/L). Based on the mortality, immobility and observation data, the 48-hour no mortaility/immobility concentration and the noobserved-effect concentration was 6.8 ug/L (nominal) (3.2 ug/L mean measured concentration).

Data Quality

Reliability

High

Data Reliability Remarks

This study was performed according to current guidelines and Good Laboratory Practices by a laboratory with considerable experience with these studies. Extensive attention was paid to analytical method development and performance.

Reference

>> Remarks

Graves, W and Swigert, J. (1997) Hexabromocyclododecane (HBCD): a 48-hour flow-through acute toxicity test with the cladoceren (Daphnia magna). Project Number: 439A-102. Wildlife International Ltd., Easton, MD.

General

EPA High Production Volume (HPV) Track Ecotoxicity End Point: Acute Toxicity to Aquatic Invertebrates

| | | <u> </u> | | |
|--------------|-----------------|---|----------------|---------|
| Sponsor ID | 1100021 | Albemarle Corporation | Create Date | 4/6, 01 |
| CAS Number | 3194556 | Cyclododecane: 1,2,5,6,9,10-hexabromo- | Study Number | 1 |
| Consortia ID | 110:012 | CMA Brominated Flame Retardant Industry Panel (BFRiP) | Completed: | N |
| | 04 - 1 | d by the Object of the Angles | 4-151 | |
| | Industry Panel. | d by the Chemical Manufacturers Association Bromina | ted Flame Reta | rdant |

Ecotoxicity End Point : Toxicity to Aquatic Plants

| Sponsor (D | 1100021 | Albemarle Corporation | Create Date | 4 18/0 1 |
|--------------|---------|---|--------------|-----------------|
| CAS Number | 3194856 | Cyclododecane, 1,2,5,6,9.10-hexabromo- | Study Number | |
| Consortia ID | 1101012 | CMA Brominated Flame Retardant Industry Panel (BFRIP) | Completed: | N |

Revision Date:

Test Substance

12/5/01

Remarks

The test article was a composite of equal parts of the commercial hexabromocyclododecane (HBCD) commercial product produced by Albemarle Corporation, Dead Sea Bromine Group, and Great Lakes Chemical Corporation. The test article composite was analyzed for characterization and homogeneity. The results of the analysis indicated the test article was homogeneous and contained the following components: HBCD beta isomer 8.5%, HBCD alpha isomer 6.0%, HBCD gamma isomer 79.1%.

Chemical Category

Method

| >> Method/Guideline folio | wed |
|-----------------------------|--|
| OECD Method 201; TSC | A Title 40, CFR, Part 797, Section 1050 |
| >> Test Type | |
| static | |
| >> GLP Yes | >> Year study performed 1997 |
| >> Species | |
| Selenastrum capricornut | JM. |
| >> End Point Cell densititi | es and area under the growth curve. |
| >> Analytical monitoring | HPLC/UV/VIS Detector; LOQ=0.400 ug/L |
| >> Exposure period | 96 Hours |
| >> Statistical Method | Shapiro Wilk's;Bartlett's;Dunnett's;Bonferroni's t |

Remarks for Method

The freshwater alga, Selenastrum capricornutum, was exposed to one of five test concentrations, a solvent control (DMF) or the negative (culture medium) control under static conditions for 96 hours. Three replicate test chambers were maintained for each treatment and control group. Nominal test concentrations were based on the solubility of the test substance in water (3.4 ug/L) and the results of an exploratory range finding toxity test. Nominal test concentrations were 1.5, 2.2, 3.2, 4.6 and 6.8 ug HBCD/L. The highest dose tested was also confirmed analytically (HPLC with UV/VIS

Ecotoxicity End Point:
Toxicity to Aquatic Plants

| Sponso | or ID | 1100021 | Albemarie Corporation | Create Date | 4.6.01 |
|--------|--------|---------|---|--------------|--------|
| CAS NE | mber | 3194556 | Cyclododecane, 1,2,5,6,9,10-hexabromo- | Study Number | |
| Consor | tia ID | 1101012 | CMA Brominated Flame Retardant Industry Panel (BFRIP) | Completed: | N |

detector).

Test solutions were inoculated with 1.0 mL of an inoculum with an approximate (~) density of 1.0 x 10E6 cells/mL to achieve a final cell density of 1.0 x 10E4 cells/mL. Samples of the test solutions were collected from each replicate test chamber at ~24 hour intervals during the test to determine cell density. Cell densities and area under the growth curve values were determined for each replicate and were used to calculate % inhibition values relative to the controls over the 96-hour exposure period. EC10, EC50 adn EC90 values were calculated, if possible, based on cell densities and area under the growth curve values for each 24-hour interval. The no-observed-effect concentration (NOEC) was determined based on statistitical evaluation of the cell densities and area under the growth curve values.

A primary stock solution was prepared by dissolving HBCD in dimethylformamide (DMF). The concentration of the stock was 0.068 mg HBCD/mL. Stock concentrations and the resultant test concentrations were prepared on a total product basis. A solvent control was prepared by diluting 250 uL DMF to 2.5 L with culture medium to yield a solvent concentration equivalent of that in the treatment groups.

Original cultures of the freshwater algae, Selenastrum capricornutum, were obtained from UTEX - The Culture Collection of Algae at the University of Texas at Austin, and have been maintained in culture medium at Wildlife International Ltd, Easton, MD. Algal cells used in this test were obtained from Wildlife International Ltd cultures that had been actively growing in culture medium for a least two weeks prior to test initiation. The control organisms were expected to exhibit exponential growth over the 96-hour exposure period. Exponential growth phase, defined as the period of growth where the algal cells are dividing at a constant rate, is indicated by the liner section of the growth curve.

The algal cells were cultured and tested in freshwater algal medium. Test chambers were sterile 250-ml Erlenmeyer flasks plugged with foam stoppers, and containing 100 mL of test or control algal medium. The test chambers were shaken continuously at 100 rpm, and held in an environmental chamber at 24 +/- 2 degrees C. Cool-white fluorescent lighting was used throughout the test (4310 +/- 431 lux). Samples of ~ 2 mL were collected from each treatment and control vessel at ~ 24 hour intervals during the 96-hour exposure. Cell counts were performed using an electronic particle counter (Coulter Electronics, Inc.). Samples of the test medium (test samples) were collected from each treatment and control group at the beginning and end of the test to measure concentrations of the test substance.

Results

| >> Nominal concentration | 0, 0.0015, 0.0022, 0.0032, 0.0046, 0.0068 mg/L | |
|---------------------------|--|--|
| >> Measured concentration | Negative control, Solvent control, and 0.0037 mg/L | |
| >> Precision > | | |

EPA High Production Volume (HPV) Track Ecotoxicity End Point: Toxicity to Aquatic Plants

| Sponsor ID | 11000 | 21 Albemarie Corpo | oration | | Create Date | 4/6/01 |
|----------------------|--------------|--|--|------------------|--------------------|-----------|
| CAS Number | 3194, | তি Cyclododocane. | 1,2,5,6,9,10-hexabromo- | | Study Number | 1 |
| Consortia ID | 11019 | 12 CMA Bronunated | Flame Retardant Industry | Panel (BFRIP) | Completed: | |
| >> Endpoint Type | EC0 | | | | | |
| >> Endpoint Value | | 0 | >> Unit used mg/L | | | |
| >> Concentration | Type Nomin | al | >> Endpoint Time | | 96 | |
| >> NOEC Precision | n > | >> NOEC | 0 | >> Unit us | ed mg/L | |
| >> NOEC Concent | ration Type | Nominal | | | | |
| >> NOEC Effect(s) | assesse | cell densisty and gro | wth | | | |
| >> LOEC Precisio | n > | >> LOEC | 0 | >> Unit use | mg/L | |
| >> LOEC Concenti | ration Type | Nominal | | | | |
| >> LOEC Effect(s) | assesse | cell density and growt | h | | | |
| >> Response of Co | ontrol Group | (was it satisfactory | ? Yes | | | |
| >> Statistical resul | ts | | | | | |
| | | | found between control a could not be defined. | nd treated grou | ps. Algal growth w | as not |
| Results Remark | | | | | | |
| | NOEC is gre | eater than HBCD's wa | n in the freswater algae ater solubility. on at the 6.8 ug/L dose le | | | he |
| Conclusions | | | | | | |
| | i | nr effect concentration ater than HBCD's wa | n in the freswater algae iter solubility. | tested could not | be determined. The | ne |
| Data Quality | Reliability | High | | | | |
| Data Reliability Ren | narks | | | | | |
| 12/20/01 | | | | | Pag | ge 3 of 7 |

EPA High Production Volume (HPV) Track Ecotoxicity End Point: Toxicity to Aquatic Plants

| Sponsor ID | 1100021 | Afbeinarle Corporation | Create Date | 4/6/01 |
|--|-------------------|---|-------------------|-----------|
| CAS Number | 3194556 | Cyclododecane, 1,2,5,6,9,10-hexabromo- | Study Number | • |
| Con sortia I D | 110:012 | CMA Brominated Flame Retardant Industry Panel (BFRIP) | Completed: | N |
| Reference | laboratory with c | performed according to current guidelines and Good Lonsiderable experience with these studies. Extensive didevelopment and performance. | | |
| >> Remarks | | I. Swigert. Hexabromocyclododecane (HBCD) A 96-Fa (Selenastrum capricornutum). Wildlife International | | |
| | | 97. Wildlife International Ltd., Easton, MD. | Lta. Froject Namo | CI. 439A- |
| (Marketon and Marketon and Mark | | | | |
| <u>General</u> | <u> </u> | | | |
| | | | | |
| | | | | |

Ecotoxicity End Point:
Toxicity to Aquatic Plants

| Sponsor ID | 110002* | Albemarle Corporation | C | reate Date | 4/6/01] |
|-------------------------------|----------------|--|---|---------------|-----------|
| CAS Number | 3194556 | Cyclododecano, 1,2,5,6,9,10-hexabromo- | S | tudy Number | 2 |
| Consortia ID | 1101012 | CMA Brominated Flame Retardant Industry | Panel (BFRIP) C | ompleted: | N |
| | | | | Revis | ion Date: |
| Test Substance | | | | | 12/5/01 |
| 1 1 | • | exabromocyclododecane (HBCD) com eat Lakes Chemicals Corporation (Wes | • | as obtained f | rom one |
| Chemical Category | ette. | | | | |
| Method | | | | | |
| >> Method/Guideline fo | llowed | | • | | |
| Not specified. | | | | | |
| >> Test Type | | | · · · · · · · · · · · · · · · · · · · | | |
| Not specified. | | | 771.771.171.771.171.171.171.171.171.171 | | |
| >> GLP Unknown | | >> Y | ear study perforr | ned 1987 | |
| >> Species | | | | | |
| Skeletonema costatun | n, Thalassions | ira pseudonana, Chlorella sp. | | | |
| >> End Point cell numb | ers | | | | |
| >> Analytical monitorin | g Capillary o | column GLC; DL not specified. | | | |
| >> Exposure period | 72 Hr S. C | ostatum, T. Pseudonana; 96 Hr Chlore | ella | | |
| >> Statistical Method | None - use | ed linear regression to determine EC50 |) | | |

Remarks for Method

Each test was replicated. Population density was estimated by cell counts on a hemacytometer. The test article was introduced into growth flasks by adding 0.05 ml test article in acetone to 51 ml growth medium with algae. Algal species tested were S. costatum (Greville) Cleve, T. pseudonana Hasle and Heindal, and Clorella sp., and were obtained from University of Rhode Island, Woods Hole Oceanographic Institution, and the Culture Collection of Algae, University of Texas at Austin, respectively. Growth media were prepared from seawater collected from an inshore site on the Gulf of Mexico and from five commercial sea salt formulations. Toxicity was expressed as the EC50 based on the cell numbers after incubation for 72 (S. costatum) or 96 hrs (T. pseudonana, Chlorella

EPA High Production Volume (HPV) Track Ecotoxicity End Point: Toxicity to Aquatic Plants

| Sponsor ID | 1100021 | Albemarle Corporation | Create Date | 4/6/01 |
|--------------|---------|---|--------------|--------|
| CAS Number | 3194556 | Cyclododecane. 1,2,5,6,9,10-hexabromo- | Study Number | 2 |
| Consortia ID | 1101012 | CMA Brominated Flame Retardant Industry Panel (BFRIP) | Completed: | N |

sp.). The EC50 was derived by straight line graphical interpolation without calculation of confidence intervals. The highest test article concentration was determined by adding the test article slowly to growth media and observing the highest concentration at which crystals did not form.

| | | | | |
|--------------------------------|--|--|--------------|--|
| Results | | | | |
| >> Nominal concentration | | | | |
| - Hollinai Concentration | | ************************************** | | |
| >> Measured concentration | | | | |
| >> Precision | | | | |
| >> Endpoint Type | | | | |
| >> Endpoint Value | 0 | >> Unit used | | |
| >> Concentration Type | | >> Endpoint Time | | 0 |
| >> NOEC Precision | >> NOEC | 0 | >> Unit used | |
| >> NOEC Concentration Type | | | | - |
| >> NOEC Effect(s) assesse | | | | |
| >> LOEC Precision | >> LOEC | 0 | >> Unit used | |
| >> LOEC Concentration Type | | | | |
| >> LOEC Effect(s) assesse | | | | Managar Harana and Araba Managar Ang and Araba Managar Ang and |
| >> Response of Control Group (| was it satisfactor | y? | - | |
| >> Statistical results | | | | |
| | Anna ta anna anna anna anna anna anna an | | | |
| | | | | |
| Results Remark | | | | |

EPA High Production Volume (HPV) Track Ecotoxicity End Point:

| Albemarle Corporation Create Date |
|---|
| Growth of Chlorella sp. was not inhibited by HBCD at the highest dose tested, 1.5 mg/L. HBCD's EC50 for S. costatum ranged from 9.0-12.0 ug/L in the six media. Similarly, HBCD's EC50 in T. pseudonana ranged from 0.05-0.37 mg/L in the six media. The pH of the six different growth media ranged from 7.6-8.2. No relationship of pH to toxicity was found for HBCD. There was little variation in the response of S. costatum to HBCD among the media, but the response to T. pseudonana varied widely. S. costatum may be more sensistive to HBCD than T. psuedonana. Conclusions HBCD's 96 hour EC50 in Chlorella sp., tested in 6 different growth media, was > 1.5 mg/L. HBCD's 72 hour EC50 in S. Costatum and T. Pseudonana in 6 different growth media ranged from 0.09-0.012 and 0.5-0.36 mg/L, respectively. All EC50 values determined in the three marine algae were greater than HBCD's water solubility (0.0034 mg/L). Reliability good |
| Growth of Chlorella sp. was not inhibited by HBCD at the highest dose tested, 1.5 mg/L. HBCD's EC50 for S. costatum ranged from 9.0-12.0 ug/L in the six media. Similarly, HBCD's EC50 in T. pseudonana ranged from 0.05-0.37 mg/L in the six media. The pH of the six different growth media ranged from 7.6-8.2. No relationship of pH to toxicity was found for HBCD. There was little variation in the response of S. costatum to HBCD among the media, but the response to T. pseudonana varied widely. S. costatum may be more sensistive to HBCD than T. psuedonana. Conclusions HBCD's 96 hour EC50 in Chlorella sp., tested in 6 different growth media, was > 1.5 mg/L. HBCD's 72 hour EC50 in S. Costatum and T. Pseudonana in 6 different growth media ranged from 0.009-0.012 and 0.5-0.36 mg/L, respectively. All EC50 values determined in the three marine algae were greater than HBCD's water solubility (0.0034 mg/L). Reliability good |
| EC50 for S. costatum ranged from 9.0-12.0 ug/L in the six media. Similarly, HBCD's EC50 in T. pseudonana ranged from 0.05-0.37 mg/L in the six media. The pH of the six different growth media ranged from 7.6-8.2. No relationship of pH to toxicity was found for HBCD. There was little variation in the response of S. costatum to HBCD among the media, but the response to T. pseudonana varied widely. S. costatum may be more sensistive to HBCD than T. psuedonana. Conclusions HBCD's 96 hour EC50 in Chlorella sp., tested in 6 different growth media, was > 1.5 mg/L. HBCD's 72 hour EC50 in S. Costatum and T. Pseudonana in 6 different growth media ranged from 0.009-0.012 and 0.5-0.36 mg/L, respectively. All EC50 values determined in the three marine algae were greater than HBCD's water solubility (0.0034 mg/L). Reliability good |
| found for HBCD. There was little variation in the response of S. costatum to HBCD among the media, but the response to T. pseudonana varied widely. S. costatum may be more sensistive to HBCD than T. psuedonana. Conclusions HBCD's 96 hour EC50 in Chlorella sp., tested in 6 different growth media, was > 1.5 mg/L. HBCD's 72 hour EC50 in S. Costatum and T. Pseudonana in 6 different growth media ranged from 0.009-0.012 and 0.5-0.36 mg/L, respectively. All EC50 values determined in the three marine algae were greater than HBCD's water solubility (0.0034 mg/L). Pata Quality Reliability good |
| response to T. pseudonana varied widely. S. costatum may be more sensistive to HBCD than T. psuedonana. Conclusions HBCD's 96 hour EC50 in Chlorella sp., tested in 6 different growth media, was > 1.5 mg/L. HBCD's 72 hour EC50 in S. Costatum and T. Pseudonana in 6 different growth media ranged from 0.009-0.012 and 0.5-0.36 mg/L, respectively. All EC50 values determined in the three marine algae were greater than HBCD's water solubility (0.0034 mg/L). Pata Quality Reliability good |
| HBCD's 96 hour EC50 in Chlorella sp., tested in 6 different growth media, was > 1.5 mg/L. HBCD's 72 hour EC50 in S. Costatum and T. Pseudonana in 6 different growth media ranged from 0.009-0.012 and 0.5-0.36 mg/L, respectively. All EC50 values determined in the three marine algae were greater than HBCD's water solubility (0.0034 mg/L). Data Quality Reliability good |
| 72 hour EC50 in S. Costatum and T. Pseudonana in 6 different growth media ranged from 0.009-0.012 and 0.5-0.36 mg/L, respectively. All EC50 values determined in the three marine algae were greater than HBCD's water solubility (0.0034 mg/L). Pata Quality Reliability good |
| |
| Data Reliability Remarks |
| |
| |
| |
| Reference |
| >> Remarks Walsh, G., Yoder, M., Mclaughlin, L., Lores, E. (1987) Responses of marine unicellular algae to brominated organic compounds in six growth media. Ecotoxicology and Environmental Safety, 14, 215-222. |
| <u>General</u> |
| |
| |

EPA High Production Volume (HPV) Track Toxicity End Point: Acute Toxicity

| Sponsor ID | 1100021 | Albemarle Corporation | | Create | Date | 37(510 |
|--------------------|--|--|--------------------------------|------------------|------------|-----------|
| CAS Number | 3194556 | Cyclododecane. 1,2,5,6 | 5,9,10-hexabromo- | Study | Number | |
| Consortia ID | 1101012 | CMA Brominated Flam | e Retardant Industry Panel (Bi | FRIP) Compl | eted: | Y |
| | | | | | Revisio | n Date: |
| | | | | | IVEAISIO | 4/10/01 |
| st Substance | | | | | | 4/10/01 |
| Remarks | A form of hexab details are avail | | HBCD) supplied as test a | rticle by Sayted | th Inc. No | o further |
| emical Category | • | | | | | |
| > Method/Guidelin | e followed | | | | | |
| Not known. | | | | | | |
| > GLP Unknown | | | >> Year stud | ly performed | 1978 | |
| > Species | | | | | | |
| rat | The state of the s | | | | | |
| > Strain no data | | | | | | |
| > Sex Both | | | | | | |
| > Number of males | s per dose | 5 | >> Number of females pe | er dose | | 5 |
| > Vehicle Corn oil | | | | | | |
| > Route of Admini | stration | | | | | |
| | | AND THE RESIDENCE OF THE PROPERTY OF THE PROPE | | | | |
| Oral | | | | | | |

| SponsorID | *************************************** | | |
|------------------|---|--|------------------|
| | 1100021 | Albemarle Corporation | Create Date 3,6/ |
| CAS Number | 3194556 | Cyclododecane: 1,2,5,6,9:10-hexabromo- | Study Number |
| Consortia ID | 1101012 | CMA Brominated Flame Retardant Industry Panel (BFRIP) | Completed: Y |
| | | roups of ten (5M:5F), 192-260 g, were administered a rved for 14 days. The highest volume used was 40 m | |
| Precision > | | | |
| Acute Lethal V | alue [| 10000 | |
| Unit mg/kg-b | | | |
| | | females died on test. | |
| One of five male | es died on test. No | females died on test. | |
| One of five male | es died on test. No | females died on test. | |
| One of five male | es died on test. No | females died on test. | |
| | s died on test. No | females died on test. | |

12/20/01

| Sponsor ID | 1100021 | Albemarle Corporation | Create Date | 4,640 |
|------------------|------------------|---|--|--|
| CAS Number | 3194556 | Cyclododecane, 1.2,5,6,9,10-hexabromo- | Study Number | |
| Consortia ID | 1101012 | CMA Brominated Flame Retardant Industry Panel (BFRIP |) Completed: | Y |
| | are consistent w | and not performed according to current guidelines. rith the general lack of toxicity associated with this n t was found acceptable. | Nonetheless, the re naterial in other mam | sults malian |
| <u>leference</u> | | | | |
| >> Remarks | | llanker, A. (1978) Final Report. Oral LD50 (Rat). E imer Product Testing Company Incorporated, Fairfic | | ∍ No.: |
| ieneral | | | | |
| | Sponsored by Sa | aytech, Inc., Sayreville, NJ. | | |
| | | | | Assessment of the second control of the seco |

| | | | V) II GOIT ADDID TO | | | |
|----------------------|---|---------------------|-------------------------------|--|------------|---------|
| Sponsor ID | (100021) A | lbemarle Corporatio | on | Creat | e Date | 1/6/01 |
| CAS Number | (19456) C | yclododecane. 1,2,5 | 5,6,9,10-hexabromo- | Study | Number [|) |
| Consortia ID | 1101012 G | MA Brominated Fla | me Retardant Industry Panel (| BFRIP) Comp | letod: | Y |
| | | | | | Davision | Detail |
| | | | | | Revision | |
| st Substance | | | | | 1 | 2/5/01 |
| | A form of hexabrom letails are available | | (HBCD) supplied as test | article by Sayte | ch Inc. No | further |
| | | | | | | · |
| nemical Category | | | | | | |
| <u>ethod</u> | | | | | | |
| > Method/Guideline | followed | | | | | |
| Not known. | | | | | | |
| > GLP Unknown | | | >> Year stu | ldy performed | 1978 | |
| > Species | | | | | | |
| rabbit | | | | | | |
| > Strain New Zeala | nd White | | | | | |
| | | | | Accessed to the second | | |
| > Sex Both | | | | | | |
| > Number of males | per dose | 3 | >> Number of females | per dose | | 3 |
| > Vehicle None | | | | | | |
| | | | | | | |
| > Route of Administ | ration | | | | | |
| Dermal | | | | g/Nation) very reg to the graph of the refugge to the state of the sta | A | |
| Remarks for Metho | d | | | | | |

| Sponsor ID | 1100021 | Albemarle Corporation | Create Date 4 6/01 |
|--|-----------------------------------|--|--|
| CAS Number | 3194556 | Cyclodedecane, 1,2,5,6,9,10-hexabromo- | Study Number |
| Consortia ID | 1101012 | CMA Brominated Flame Retardant Industry Panel (BFRIP | Completed Y |
| | half with abrade application dern | scribed as that of Hagen (1959). Albino rabbits in great skin, 1.88-2.07 kg, highest dose level mechanicall nally under occluded patch, observed for 14 days. In the sible due to mechanical and physical limitations is 8 states. | y possible, single Material used as received. |
| Results >> Precision > | | | |
| >>Acute Lethal Val | ue | 8000 | |
| >> Unit mg/kg-bw >> Deaths per Dos | | | |
| No animals died or | | | |
| Results Remark | | | |
| | | | |
| Conclusions | The domest DE | O of UPCD in makita was > 0.000 mar/km hadu waiah | |
| | me dermar LDS | 0 of HBCD in rabbits was > 8,000 mg/kg body weigh | IL . |
| Data Quality | Reliability Acc | peptable. | |

Data Reliability Remarks

| Sponsor (D | | | | |
|--------------|--|---|--------------|-------------|
| | 1100021 | Albemarle Corporation | Create Date | 4.0 |
| CAS Number | 3194556 | Cyclododecane. 1.2.5,6.9,10-hexabromo- | Study Number | |
| Consortia ID | 1101012 | CMA Brominated Flame Retardant Industry Panel (BFRIP) | Completed: | Y |
| | are consistent w | l and not performed according to current guidelines. vith the general lack of toxicity associated with this m t was found acceptable. | | |
| | | | | |
| Reference | | | | |
| >> Remarks | | alanker, A. (1978) Final Report. Dermal LD50 (Rabl Consumer Product Testing Company Incorporated, F | | eference |
| | | | | |
| | | | | |
| Seneral | | | | |
| | Anna and a superior a | | | *** 181 781 |

| | | | - / | <u> </u> | | |
|--|--------------------------------------|----------------------------|-----------------------|-------------------|--|------------|
| Spensor ID | 1100021 | Albemarle Corporation | on | | Create Date | 1/6/01 |
| CAS Number | 3194556 | Cyclododecane, 1,2,5 | 5.6,9,10-hexabromo- | | Study Number | 3 |
| Consortia ID | 1101912 | CMA Brominated Fla | me Retardant Industry | Panel (BFRIP) | Completed: | Υ |
| | | | | | Revisi | on Date: |
| | | | | | | 4/11/01 |
| <u> [est Substance</u> | | | | | *************************************** | |
| | A form of hexab details are avail | romocyclododecane able. | (HBCD) supplied a | s test article by | Saytech Inc. N | No further |
| TO COLUMN AND ADDRESS OF THE ADDRESS | | | | | | |
| | | | | | No World and an additional and additional additional additional and additional ad | |
| Chemical Category | | | | | | |
| Method | | | | | | |
| >> Method/Guideline | followed | | | | | |
| Not known. | | | | | | |
| >> GLP Unknown | | | >> Ye | ear study perfo | rmed 1978 | |
| >> Species | | | | | | |
| rat | | | | | | |
| >> Strain no data | | | | | | |
| >> Sex Both | | | | | | |
| >> Number of males | per dose | 5 | >> Number of fem | nales per dose | | 5 |
| >> Vehicle None | | | | | | |
| >> Route of Adminis | tration | | | | | |
| Inhalation | | | | | | |
| Remarks for Meth | od | 4 - | | | | |

| | 1100621 | Albemarle Corporation | Create Date 1/3 |
|-----------------|--------------|--|-----------------|
| CAS Number | 319 (556) | Cyclododecane, 1,2,5,6,9,10-hexabromo- | Study Number |
| Consortia ID | 1101012 | CMA Brominated Flame Retardant Industry Panel (BFRIP) | Completed: Y |
| | | oups of 10 (5M:5F), 233-292 g, exposed to concentrati er concentration) for one hour, observed two weeks. M | |
| sults Precision | | | |
| Acute Lethal \ | Value | 200 | |
| • Unit mg/L(a | ir) | | |
| Deaths per D | lose | | |
| No animals died | d on test. | | |
| | | | <i>1</i> |
| Results Remai | rk | | |
| Results Remai | rk | | |
| Results Remai | rk | | |
| | | | |
| Results Remai | | C50 of HBCD in rats was > 200 mg/L for a 1 hour expo | |

| Spensor ID | 100021 | Albemarle Corporation | Create Date | |
|--------------|--|---|----------------------|----------------|
| CAS Number | 3194556 | Cyclododecane. 1,2,5,6,9.10-hexabromo- | Study Number | |
| Consortia ID | 1101012 | CMA Brominated Flame Retardant Industry Panel (BFRIP) | Completed: | Y |
| | are consistent v | d and not performed according to current guidelines. with the general lack of toxicity associated with this mit was found acceptable. | Nonetheless, the rea | sults malia |
| | | | | |
| erence | Section and the section of the secti | | | |
| | | alanker, A. (1978) Final Report. Inhalation LC50 (Ra Consumer Product Testing Company Incorporated, F | | rence |
| | | | | erenci |
| Remarks | | | | renco |
| >> Remarks | No.: 78385-2. C | | | erer |

EPA High Production Volume (HPV) Track Toxicity End Point: Developmental Toxicity/Teratogenicity

| Sponsor ID | [10002] | Albemarie Corporation | Create Date | 4,6/01 |
|---------------------------|--|---|--|---|
| CAS Number | 310.1956 | Cyclododecane, 1,2,5,6,9 10-hexabromo- | Study Number | 1 |
| Consortia ID | 1101012 | CMA Brominated Flame Retardant Industry Panel (BFRIP) | Completed: | |
| | | | | |
| | | | Revision | |
| est Substance | | | 1 | 2/5/01 |
| Remarks | commercial prod Chemical Corpor homogeneity. T | vas a composite of equal parts of the commercial hex luct produced by Albemarle Corporation, Dead Sea B ration. The test article composite was analyzed for cl he results of the analysis indicated the test article was nponents: HBCD beta isomer 8.5%, HBCD alpha isor | romine Group, and naracterization and s homogeneous an | Great Lakes d contained |
| hemical Category | | | | |
| ethod >> Method | d/Guideline follo | wed | | |
| EPA OPPTS Me | thod 870.3700; OI | ECD 414 | | |
| >> GLP Yes | | >> Year study pe | rformed 1999 | |
| >> Species | _ | | | |
| rat | | | | |
| | I l strai Sprague- | Dawley | | |
| >> Sex F | 1 | | | |
| >> Number of male | s per dose | 0 >> Number of females per dos | e | 25 |
| >> Route of Admin | istration Oral | | | |
| >> Days of Gestation | | | | |
| | | | | |
| >> Frequency of tre | eatment Once | daily | | |
| | | | | |
| >> Doses 0, 250, 5 | 500, 1000 mg/kg b | pody weight | | |
| >> Control Group | Yes | Concurrent control | | |
| >> Statistical Metho | | | | |
| See Remarks for | | | | |
| | | | | |
| Remarks for Met | hod | | | parties and the second |
| | · · · · · · · · · · · · · · · · · · · | | | |

EPA High Production Volume (HPV) Track

Toxicity End Point:
Developmental Toxicity/Teratogenicity

| Sponsor ID | 1100021 | Albemarle Corporation | Create Date | 4/6/01 |
|--------------|---------|---|--------------|--------|
| CAS Number | 3194556 | Cyclododecane. 1,2.5,6.9,10-hexabromo- | Study Number | 1 |
| Consortia (D | 1101012 | CMA Brominated Flame Retardant Industry Panel (BFRIP) | Completed: | |

Hexabromocyclododecane (HBCD) was administered by gavage in corn oil to three groups of 25 bred Crl:CD(SD)IGS BR (Charles River Laboratories, Raleigh, NC) rats once daily from gestation days 6 through 19. Dosage levels were 250, 500 and 1000 mg/kg/day administered in a dose volume of 5 ml/kg. A concurrent control group composed of 25 bred females received the vehicle, corn oil, on a comparable regimen. Clinical observations, body weights and food consumption were recorded. On gestation day 20, a laparohysterectomy was performed on all animals. The uteri and ovaries were examined and the numbers of fetuses, early and late resorptions, total implantations and corpora lutea were recorded. Mean gravid uterine weights and net body weight changes were calculated for each group. The fetuses were weighed, sexed and examined for external soft tissue and skeletal malformations and variations.

Appropriate statistical tests were used for each end point and included a one-way ANOVA with Dunnet's test, and Kruskal Wallis test with Mann-Whitney U test.

Results

| >> Maternal Precision/NOAEL | = | | | | |
|---------------------------------|-------------|-----------|--------------|----------|---|
| >> Maternal NOAEL dose | 1000 | | >> Unit used | mg/kg-bw | |
| >> Maternal NOAEL effect None |) | | | | |
| >> Maternal Precision/LOAEL | > | | | | |
| >> Maternal LOAEL dose | 1000 | | >> Unit used | mg/kg-bw | |
| >> Maternal LOAEL effect None |) | | | | |
| >> Developmental Precision/NO | AEL = | | | | |
| >> Developmental NOAEL dose | | 1000 | >> Unit used | mg/kg-bw | |
| >> Developmental NOAEL effect | None | | | | |
| >> Developmental Precision/NO | AEL > | | • | | |
| >> Developmental LOAEL dose | | 1000 | >> Unit used | mg/kg-bw | |
| >> Developmental LOAEL effect | None | | | | |
| >> Actual dose | | | | | |
| As given above. | | | | | |
| >> Maternal data with dose leve | l (with NOA | EL value) |)• | | |
| | | | | | |
| No adverse effects detected. | | | | | 7999991100 Block-land and a second a second and a second a |
| , | | | | | |
| | | | | | |

EPA High Production Volume (HPV) Track Toxicity End Point: Developmental Toxicity/Teratogenicity

| Sponsor tD | 110002 | 2t Alb | oemarle Corporat | tion | | Create Date | -1/6/01 |
|----------------------|---|--|--|---|--|--|----------------------------|
| CAS Number | 319455 | 56 Cyc | clododecane. 1.2 | 2.5,6,9,10-hexabro | emo- | Study Number | 1 |
| Consortia ID | 110101 | 12 CM | IA Brominated FI | lame Retardant Ir | idustry Panel (BFRIP | Completed: | |
| >> Fetal data with | dose level (v | with NO | AEL value). | | | | |
| No adverse dete | ected. | *************************************** | | | | | |
| >> Statistical resu | ilts | | | | | | |
| See Methods. | | on description of the statement when the statement of the | | | | | |
| Results Remark | *** | | ann a' bhail ann ann an ann an an Airth Madrin ann ann an airth ann ann ann ann ann ann ann ann ann an | | | THE RESEARCH STATE OF THE STATE | |
| Conclusions | laparohyster Body weight necropsy, no unaffected b | rectomy. t gain and o treatme by test art | No treatment- d food consument-related find ticle administra | -related clinica option were not lings were obso ation at any do | signs were obser adversely affected erved. Intrauterine se level. No treatr | amined at the sche rved at any dose le d at any dose level e growth and survive ment-related fetal the treated groups | evel. I. At val were |
| | | | | vel for materna days 6-19 of go | | lopmental toxicity | was 1000 |
| Data Quality | Reliability | High | | | | | |
| Data Reliability Ren | narks | | | | | | |
| | | | | | | ed dose studies ur rmance of studies | |
| Reference | · · | | | | | | |
| >> Remarks | | Rats. Lab | | | | bromocyclododeca n Laboratories, Inc | |
| <u>General</u> | | - 10 to 1 t | | | | | |
| | Sponsored by Panel. | by Chemic | cal Manufactur | rers Associatio | n Brominated Flan | ne Retardant Indus | stry |

EPA High Production Volume (HPV) Track

Toxicity End Point:
Developmental Toxicity/Teratogenicity

| LIA Ingil I roduction volume (III v) I ruck Devek | pmental loxicity/leratogenicity |
|--|--|
| Spensor 4D 1100021 Albemarle Corporation | Create Date |
| CAS Number 3194556 Cyclododecane, 1,2,5,6,9.10-hexabromo- | Study Number |
| Consortia ID 1101012 CMA Brominated Flame Retardant Industry Pane | (BFRIP) Completed: |
| | Revision Date: |
| est Substance | 12/5/01 |
| Remarks The test article was manufactured by Daiichi Kogyo Seiyaku composition is known. | K.K. No further information on its |
| hemical Category | |
| ethod >> Method/Guideline followed | |
| Not specified. | THE RESERVE OF THE PROPERTY OF |
| >> GLP Unknown >> Year | study performed 1985 |
| >> Species | |
| rat | |
| >> Strain Mammal strai Wistar | |
| >> Sex F | |
| >> Number of males per dose 0 >> Number of females | per dose 20 |
| >> Route of Administration Oral | |
| >> Days of Gestation 0-20 | |
| | |
| >> Frequency of treatment Daily | |
| | |
| >> Doses 0, 0.01, 0.1 and 1% of the Diet | |
| >> Control Group Yes Concurrent control | |
| >> Statistical Method | |
| Not specified. | |
| | |
| Remarks for Method | |

EPA High Production Volume (HPV) Track Developmental Toxicity/Teratogenicity

Toxicity End Point:

| Sponsor ⁽ D | 1100021 | Albemarle Corporation | Create Date | 4/6/01 |
|------------------------|---------|---|--------------|--------|
| CAS Number | 3194556 | Cyclododecane: 1,2,5.6,9.10-hexabromo- | Study Number | 2 |
| Consortia ID | 1101312 | CMA Brominated Flame Retardant Industry Panel (BFRIP) | Completed: | |

The Murai et al study consisted of a 7 day dose range finding study (n=5 rats/dose group) and a combined teratogenicity-developmental study (n=20/dose group). Doses in the 7 day range finding study were 0, 0.3, 1, 3 or 10 g/kg/day. Doses as high as 10 g/kg/day produced no evidence of toxicity. A statistically significant (P<0.01) increase in liver weight was noted in groups receiving > 1 g/kg/day. Doses for the combined teratogenicity-developmental study were based on this increase in liver weight.

In the combined teratogenicity-developmental study, pregnant female rats were fed diets containing 0, 0.01, 0.1, or 1% HBCD on days 0-20 of gestation. Daily doses were estimated by the authors to be 0, 5, 50 or 500 mg/kg/day and the average total dose/rat/group was estimated to be 0, 0.13, 1.28 or 12.0 g/kg. Rats were observed daily and body weight and food consumption measured. Fourteen rats from each group were sacrificed on day 20 of gestation and their fetuses were examined for toxicity or teratogenicity. Approximately 150 fetuses/dose level were examined for evidence of teratogenicity. All fetuses from all litters were examined for signs of external anomalies. Approximately 2/3 of the fetuses/dam were examined for skeletal abnormalities; the remaining fetuses from each dam were examined for any abnormalities of the internal organs. In addition, six rats from each group were allowed to deliver their litters and growth of the litters was observed until the 7th week post-parturition.

Results

| >> Maternal Precision/NOAEL | > | | |
|--------------------------------|---------------------|---------------------------------------|---|
| >> Maternal NOAEL dose 1000 | | >> Unit used | mg/kg in feed |
| >> Maternal NOAEL effect No | adverse effects, in | ncreased liver wt at 1% dose. | |
| >> Maternal Precision/LOAEL | > | | |
| >> Maternal LOAEL dose | 1000 | >> Unit used | mg/kg in feed |
| >> Maternal LOAEL effect No | adverse effecs. | | |
| >> Developmental Precision/N | OAEL > | | |
| >> Developmental NOAEL dos | e 100 | 00 >> Unit used | mg/kg in feed |
| >> Developmental NOAEL effe | ct No adverse ef | fects. | |
| >> Developmental Precision/N | OAEL > | · · · · · · · · · · · · · · · · · · · | |
| >> Developmental LOAEL dos | 100 | 0 >> Unit used | mg/kg in feed |
| >> Developmental LOAEL effe | ct No adverse eff | ects. | |
| >> Actual dose | | | |
| Estimated as 0, 5, 50, 500 mg | HBCD /kg bd wt/d | lay | 11 Marie 11 |
| >> Maternal data with dose lev | rel (with NOAEL) | value). | agan pagan saran af |

EPA High Production Volume (HPV) Track

Toxicity End Point:
Developmental Toxicity/Teratogenicity

| Sponsor ID | (160021) | Albertarie Corporation | Create Date | 1:6/01 |
|-------------------|---|---|--|--------|
| CAS Number | 3194506 | Cyclododecane: 1,2,5,6,9,10-hexabromo- | Study Number | 2 |
| Consortia ID | 1103612 | CMA Brominated Flame Retardant Industry Panel (BFRIP) | Completed: | |
| Only effect de | etected in dams was | an increase in liver weight at the 1% dose level. | | |
| | Market Market Market State (Market Market | | THERE IS NO THE PARTY OF THE PA | |
| >> Fetal data w | ith dose level (with | NOAEL value). | | |
| No effects de | tected in fetuses. | | | |
| | | | | |
| >> Statistical re | sults | | | |
| See results re | emarks. | | n natura papaten. Paran ini mata Majala jala ini harbitan mendada, bermata dan menen | |
| Results Rema | rk | | | |

The authors' estimated the doses in the feed were equivalent to 0, 5, 50 or 500 mg HBCD /kg body weight /day. No adverse effects were detected in any treatment group with respect to maternal weight gain, food consumption, or gross apprearance of internal organs. The mean liver (absolute and relative to body weight) weight in the 1% group was statistically different (higher) from the control mean. Normal development was seen in neonates carried through to six weeks of age.

There was no adverse effect of treatment on the number of corpera lutea, implants, resporptions, live fetuses, sex ratio, or body or placental weight. No fetal deaths occured in any group. No external, skeletal or viseral malformations were detected. A few skeletal variations were detected but where of similar types and numbers in the control and treated groups.

There was no significant differences between the control and treated groups in the number of implantation, live newborns, dead newborns, live newborn parturition index. The weaning and survival index was comparable in the control and treated groups. Body weight chnages in the newborns was comparable in all groups.

<u>Conclusions</u>

No reproductive or developmental effects where detected in rats at HBCD doses up to 1% in the diet (~500 mg/kg/d) administered from days 0-20 of gestation. Further, normal development was seen in neonates carried through to six weeks of age.

Dose levels: 0, 0.01, 0.1, or 1% HBCD on days 0-20 of gestation [Murai estimate: 0, 5, 50 or 500 mg/kg/day]. No teratogenic effects. Normal development in neonates carried through age 6 wks. NOEL = 1% of diet.

| Data Quality | | | | |
|---------------------|-------------|----|-----|----|
| | Data | Qu | ali | ty |

Reliability

Good.

Data Reliability Remarks

EPA High Production Volume (HPV) Track Toxicity End Point: Developmental Toxicity/Teratogenicity

| Sponsor ID | 1100021 | Albemarle Corporation | Create Date | 4/6/01 |
|----------------|--|--|----------------------|-------------|
| CAS Number | <i>4</i> 79.2456 | Cyclododecane. 1,2,5,6,9.10-hexabromo- | Study Number | 2] |
| Consortia ID | 1101012 | CMA Bronunated Flame Retardant Industry Panel (BFRIP) | Completed: | |
| | One author of the branch. | nis study was associated with the National Institute of I | Hygienic Science, Os | saka |
| Reference | Landania da decembra da persona d | | | |
| >> Remarks | 1 | saki, H., Kanoh, S. 1985. Studies on the tosicity of ins gnant rats - Fetal toxicity of Hexabromocyclododecane 31-986. | | |
| <u>General</u> | | | | |
| | Funding for this | study was provided by Japan's Ministry of Health and | Welfare. | |

| Sponsor ID | 1100021 Albei | marle Corporation | | Create Date | 4.6/01 |
|--|---|---|--|--|--|
| CAS Number | 019 (00a) Cycl e | ododecane, 1.2.5.6,9.10-hex | cabromo- | Study Number | ; ; |
| Consortia ID | 1101012 CMA | Brominated Flame Retarda | int Industry Panel (BFRIP) | Completed: | - 10 mg - 10 m |
| | Physical | consequences (Consequences) Società de graphy a vincia (Consequences) | | Revisio | on Date: |
| Test Substance | | | | | 12/5/01 |
| | commercial product pr Lakes Chemical Corpo homogeneity. The res | oduced by Albemarle Coration. The test article outs of the analysis indi | s of the commercial hexal corporation, Dead Sea Bro composite was analyzed cated the test article was 8.5%, HBCD alpha isome | omine Group, ar for characteriza homogeneous a | nd Great ation and and contained |
| Chemical Category Method | | | | | |
| >> Method/Guidelir | ne followed | | | | |
| OECD Method 4 | 07 | | | | |
| >> GLP Yes Were Good Labora >> Species | tory Practices followe | d in the st | >> Year study per | formed 1997 | , |
| rat | | - | | | |
| >> Strain Mamma | Sprague-Dawle | еу | | | |
| >> Sex Both >> Number of male | s per dose | 6 >> Num | ber of females per dose | | 6 |

| >> Route of Administration | on Oral | | | |
|---|--|--|--|--|
| >> Exposure Period | 28 | | | |
| Duration of study in days | s (for example, 28 days, 90 d | | | |
| >> Frequency of treatme | ent Once per day | | | |
| Number of doses per day | y, week, etc. This is particularly relevant for inhalation experiments 6hrs/day, 5 d | | | |
| >>Doses 0, 125, 350, 10 | 00 mg/kg/day; dosage volume=5 ml/kg | | | |
| List all doses used in te | | | | |
| >> Control Group Yes | | | | |
| Concurrent contro | | | | |
| >> Post observation period 14 Days | | | | |
| Length of time animals of | observed after last d | | | |
| >> Statistical Method See Remarks for Method section. | | | | |

Remarks for Method

Cite statistical methods use

Hexabromocyclododecane (HBCD) was administed orally by gavage in corn oil to three groups of Sprague-Dawley Crl:CD BR (Charles River Laboratories, Inc., Portage, MI) rats for a period of 28 consecutive days at doses of 125, 350 or 1000 mg/kg/day administered in a dosage volume of 5 ml/kg. The test groups consisted of 6 males and 6 females in the 125 and 350 mg/kg/day groups, and 12 males and 12 females in the 1000 mg/kg/day group. A concurrent control group (n=12 males and females) was treated in a similar manner with the vehicle, corn oil. At the end of the dosing period, 6 animals/sex/group were euthanized and necropsied. The remaining 6 animals/sex in the control and 1000 mg/kg/day groups remained on test untreated for a 14 day recovery period. At the end of the recovery peroid, all animals were euthanized and necropsied. Animals were 6 weeks of age at study initiation.

Animals were observed twice daily for mortality and morbundity. Clinical signs were recorded daily. Body weights and food consumption were measured weekly. Functional observational battery and motor activity evaluations were performed during weeks 1 (pretest), 3, and 5 (recovery). Samples for hematology and serum chemistry evaluations were collected at the primary (28 day) and recovery (42 day) necropsies. Complete necropsies were performed on all rats. The brain, liver, kidney, heart, spleen, testes and epidymus or ovaries, adrenal glands, and thymus from all animals were weighted at each necropsy. Approximately 40 tissues were

Results

collected and preserved at each necropsy from each animal. The following tissues were examined microscopically from the control and high dose animals: liver, kidney, heart, spleen, testes (males), prostate (males), seminal vesices (males), epididymus (males), ovaries (females), adrenal glands, thymus, bone with marrrow (sternebra), brain, stomach, cecum, duodenum, ileum, jejunum, lymph node, peripheral nerve (sciatic), spinal cord, lung, trachea, uterus (females), urinary bladder, and all gross lesions. The lungs, liver, kidney, stomach, gross lesions and target organs were examined in all dose levels.

Body weights, weight gain, food consumption, functional observation battery and motor activity results of treated animals were compared statistically by sex and treatment day to their respective control groups (p<0.05 or <0.01).

Concentrations of the dosing suspensions were confirmed. Homogeneity determinations were performed on study days 0, 13, and 27.

All statistical analyses were conducted using two-tailed tests for minimum significance levels of 1% and 5% comparing the treatment groups to the vehicle control group by sex. Analysis of body weight change, food consumption, clinical pathology values, continuous functional observational battery data and absolute and relative organ weight data were analyzed with a one-way analysis of variance followed by Dunntt's test. Discontinuous (ordinal or descriptive) functional observational battery data were analyzed using Fisher's exact test. Statistical tests for locomotor activity data were performed using SAS/STAT statistical software. Clinical laboratory values for cell types that occur at a low incidence (i.e., monocytes, eosinophils and basophils) were not subjected to statistical analysis.

| >> NOAEL Precision | >= | |
|--------------------------------------|--|---|
| >> NOAEL dose | 1000 >> Unit mg/kg-bw | |
| >> NOAEL Effec | Increase in liver weight in the absence of his | topathlogic or clincal chemistry changes. |
| (e.g., decrease in bod weight, organ | y | |
| >> LOAEL Precision | > | |
| >> LOAEL dose | 1000 >> Unit mg/kg-bw | |

| >> LOAEL Effect (e.g., decrease in be weight, organ | 1 | ne noted. | | |
|---|----------------|--|--|--|
| >> Actual dose rece | ived by dose | level by sex (if availabl | | |
| Test article admini | stered by gava | ge. | | |
| >> Toxic response | A brief narra | f narrative describing toxic response or effects, by dos | | |
| | No evidence o | f toxicity was observed at any dose level. | | |
| >> Statistical result | s Note statis | stical results, with appropri $ ho$ value | | |
| See Results Rema | rks section. | | | |

Results Remark Provide at a minimum qualitative descriptions of elements where dose effect related observations of elements were seen:

Survival was not affected by administration of the test article. All animals survived to the scheduled necropsy. Clinical signs observed during the study were nonspecific, low in incidence, non-dose-related, and not considered related to test article.

Body weights, weight gain and food consumption were not affected by treatment. No statistically significant differences in mean body weight between control and treated animals were detected with the exceptoin of an increase in mean femal body weight in the 350 mg/kg/day group during week 2. Mean female body weight at that time point was 196 g in the 350 mg/kg/day group vs. 179 g in the control group. No statistically significant differences in body weight gain between the control and treated animals with the expectation of a decrease in mean male body weight gain in the 1000 mg/kg/day recovery group during week 1 of recovery. Mean male body weight gain at that time point was 21 g vs 31 g in the control group; mean male body weight was not statistically different from the control mean. No statistically significant differences in food consumption between control and treated animals were detected with the exception of an increase in mean female food consumption in the 350 mg/kg/day during weeks, -1, 1, and 2 of treatment. Mean female food consumption at those time points were 18, 17 and 17 g vs. 16, 15 and 15 g in the control group, respectively.

Results of the functional observation battery and motor activity tests were not affected by treatment. No statistically significant differences were observed between the control and

treated animals at any time point (p<0.05).

No statistically significant differences between control and treated animals were found for hematology parameters with the exception of an increase in mean activated partial thromboplastin time in the 1000 mg/kg/day males on week 4 and a decrease in the mean prothrombin time in the 1000 mg/kg/day females on week 4. These statistical differences were not of toxicological significance.

No toxicologically significant effects on serum chemistry values related to test article administration were observed at the 28 day primary and 42 day recovery necropsies. Scattered instances of statistically significant differences between treated and control animals were detected for some serum chemistry parameters at the 28 day primary necropsy. These scattered statistical differences were not considered toxicologically significant because the statistical differences occurred in the absence of a dose response, in the absence of the accompanying clinical chemistry changes expected, in the opposite direction from what occurs in a toxic stae, in a direction which is without physiologic significance, or due to potential interference with the laboratory method. No statistically significant differences in serum chemistry parameters were detected between groups at the 42 day recovery necropsy.

No gross lesions attributable to test article administration were detected at either necropsy. Gross lesions were nonspecific, low in incidence, non-dose-related, and considered incidental.

No microscopic lesions attributable to test article administration were detected on histopatholgic exam. Microscopic changes were nonspecific, low in incidence, non-dose-related and considered incidental.

No statistically significant differences in organ weight or organ to body weight ratios were detected between control and treated animals with one exception. Absolute liver weights were statistically significantly increased with respect to control mean at the 28 day necropsy in males in the 1000 mg/kg/day group and in females in the 350 and 1000 mg/kg/day groups. Liver to body weight ratios in the 350 and 1000 mg/kg day male and female groups were statistically increased at the 28 day necropsy. At the recovery necropsy, male absolute and liver to body weight ratio were statistically comparable to the contol mean. Female absolute liver weight and liver to body weight ratio were statistically increased compared to the control mean. The difference in absolute liver weight between control and treated females was less pronounced at the end of the recovery period, indicating the increase in liver weight was reversible in females as well as males. In the absence of test article related histologic and serum chemistry changes, increases in liver weight are considered an adaptive rather than toxic response, are not uncommon in the rat, and are most likely the result microsomal

| 117 | 7 | uction |
|-----|---|-----------------|
| | u | <i>1</i> UIUI I |

Conclusions

Input optional information or further explain the contents of a particular section, much as is done in the "Discussion" portion of a publication in academic journals.

No systemic toxicity was observed at any dose level. The No Observed Adverse Effect Level of HBCD administered orally to male and female rats for 28 consecutive days was > or = 1000 mg/kg/day, the highest dose tested.

Data Quality

Reliability High

Denote the reliability of data, at the discretion of the person preparing the robust su

Data Reliability Remarks Add comments about how reliability of data was determined, or add re

This study was performed according to current guidelines for repeated dose studies under Good Laboratory Practices by a laboratory experienced in the performance of studies of this type.

Reference

Cite the full reference for the critical study on which the robust study summary is based. List other appropriate references that support this summary and, have been reviewed for

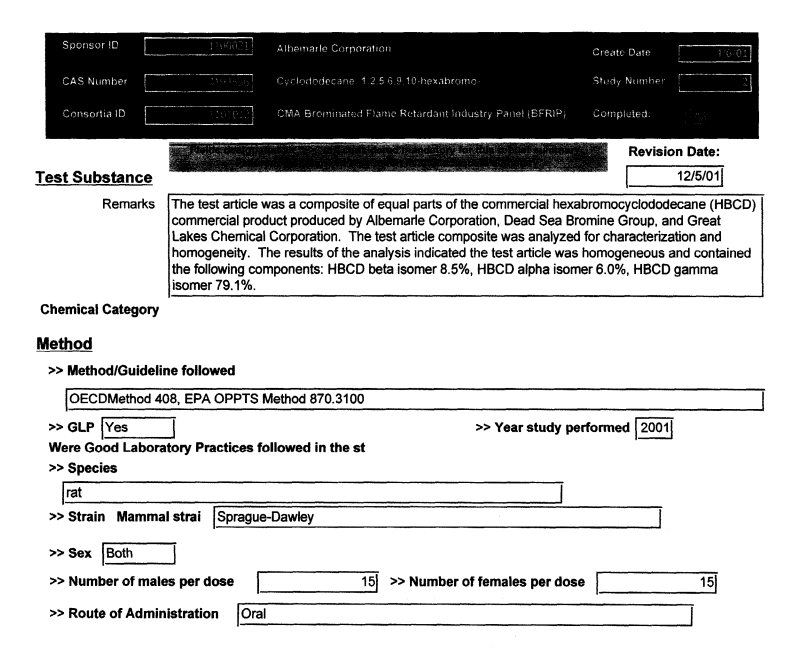
>> Remarks

Chengelis, C. (1997) A 28-Day Repeated Dose Oral Toxicity Study of HBCD in Rats. Laboratory Study Number: WIL-186004. WIL Research Laboratories, Inc., Ashland, OH.

General

Add any statement that doesn't fit into any of the other f

Study sponsored by the Chemical Manufacturers Association Brominated Flame Retardant Industry Panel.



| >> Exposure Period | 90 | | | | | |
|---|---|--|--|--|--|--|
| Duration of study in day | rs (for example, 28 days, 90 d | | | | | |
| >> Frequency of treatm | >> Frequency of treatment Once daily by gavage | | | | | |
| Number of doses per da | ny, week, etc. This is particularly relevant for inhalation experiments 6hrs/day, 5 d | | | | | |
| >> Doses 0, 100, 300, 10 | 000 mg/kg-bw; dose volume = 5 ml/kg | | | | | |
| List all doses used in te | | | | | | |
| >> Control Group Yes | | | | | | |
| Concurrent contro | | | | | | |
| >> Post observation period 30 day recovery period | | | | | | |
| Length of time animals observed after last d | | | | | | |
| >> Statistical Method | ANOVA, Dunnett's test, Others | | | | | |

Remarks for Method

The test article, a composite of three lots of commercial hexabromocyclododecane (HBCD), was administered by oral gavage in corn oil once daily to four groups of CrI:CD(SD)IGS BR rats (n=15/sex/group) at dose levels of 0 (control), 100 (low), 300 (mid) and 1000 (high) mg/kg/day seven days per week for 90 days. The dosage volume was 5 ml/kg. The control animals received the vehicle, corn oil, only. At the end of the 90-day treatment period, 10 animals/sex/group were euthanized and necropsied. The remaining rats continued on test untreated for a 28-day recovery period prior to necropsy.

Results

Cite statistical methods use

In addition to the main toxicology groups, two satellite groups of 20 animals/sex/group were treated concurrently in an identical manner at dose levels of 0 or 1000 mg HBCD/kg/day for up to 90 days. Body weights were recorded weekly. Two animals/sex/group were euthanized on study days 2, 6, 9, 13, 20, 27, 55, 89, 104 and 118, and blood and body fat (mesenteric and/or omental) were collected. The body fat was analyzed for HBCD content.

Animals in the main toxicology groups were observed twice daily throughout the study for mortality and morbidity. Body weights and food consumption were measured weekly. Blood was collected at study weeks 3 (n=5/sex/group), 13 (n=10/sex/group) and 17 (n=5/sex/group) for hematology, serum chemistry and hormone (T3, T4 and TSH) measurements. Urine was collected prior to each necropsy, at study weeks 13 and 17, for urinalysis. Ocular examinations were performed prior to study initiation and during study weeks 12 and 15.

Functional Observational Battery and Locomotor Activity evaluations were performed on 5 animals/sex/group prior to study initiation, during the last week of test article administration (study week 13), and during the recovery period. An examination of vaginal cytology (for estrus cycle determinations) was performed on study days 69-90. At each necropsy, sperm motility/viability, morphology, and number were assessed. Complete necropsies were performed on all animals. Approximately 40 organs or tissues/animal were collected and preserved. The adrenals, brain, epididymides, heart, kidneys, liver, ovaries, prostate, spleen, testes, thymus, thyroids with parathyroids, and uterus with cervix were weighed. Paraffin sections of tissues stained with hematoxylin and eosin from the control and 1000 mg/kg/day dose groups and the liver, lungs and thyroid glands in the 100 and 300 mg/kg/day doses, and gross lesions from all animals were examined under the light microscope. Livers from five randomly chosen animals/sex from the control and 1000 mg/kg/day dose groups were examined microscopically using Oil Red O or periodic acid Schiff's (PAS) reagent for evidence of lipid accumulation or glycogen accumulation/depletion, respectively. Statistical comparisons by sex and treatment day were made between the control and treated animals where indicated (p<0.05).

| >> NOAEL Precision | >= |
|---|---------------------------------------|
| >> NOAEL dose | 1000 >> Unit mg/kg-bw |
| >> NOAEL Effec | See Results Remarks. |
| (e.g., decrease in body weight, organ | ly |
| >> LOAEL Precision | > |
| >> LOAEL dose | 1000 >> Unit mg/kg-bw |
| >> LOAEL Effect (e.g., decrease in body weight, organ | No adverse effects detected. |
| | ved by dose level by sex (if availabl |
| As given under Dose | es. |

>> Toxic response A brief narrative describing toxic response or effects, by dos

| See Results Remarks. | | | |
|----------------------|--|--|--|
| | | | |

>> Statistical results ρ Note statistical results, with appropri ρ value

See Results Remarks.

Results Remark Provide at a minimum qualitative descriptions of elements where dose effect related observations of elements were seen:

No test article-related effect on mortality occurred. Clinical signs were non-specific, low in incidence, non-dose-related and not related to test article administration. No test article-related changes occurred in body weight, food consumption, Functional Observational Battery or Locomotor Activity. No test article-related effects on hematologic parameters were noted. No test article?related ocular lesions were detected at the ophthalmic exams. No test article-related changes were noted on the estrus cycle as determined by vaginal cytology, or on sperm motility/viability, morphology, and number. Instances of statistically significant differences between control and some treatment groups were detected at study week 13 in the clinical chemistry data, hormone data, organ weight data and histology findings. They were generally secondary to the inducing effects on the liver or were otherwise not considered adverse effects of treatment as discussed further below.

Statistically significant (p<0.05) test article-related clinical chemistry changes at week 13 include an increase in albumin (all dose levels for males), total protein (all dose levels for females and 1000 mg/kg/day for males), globulin (300 and 1000 mg/kg/day for females), and chloride (all doses for both sexes). In addition, increased gamma glutamyltransferase levels were noted in the 1000 mg/kg/day group (p<0.05). Thyroxine (T4) levels were decreased at study week 13 compared to the control mean in all male dose groups and the 300 and 1000 mg/kg/day dose females (p<0.05). There were no corresponding statistical effects on T3 and TSH. While potentially test article-related, the changes in serum chemistry parameters were not of sufficient magnitude to be adverse, occurred in otherwise clinically normal animals, tended to be within or close to historical control values, and were not present at the end of the recovery period; furthermore, these serum albumin and gamma glutamyltransferase increases were probably secondary to the increases in liver weight. The increases in serum chloride were probably secondary to be presence of free bromide in the test article preparation which interfered with the chloride determination methodology. The decrease in T4, which was also reversible, was also probably secondary to increased liver weight (secondary to microsomal enzyme induction, known to cause increased metabolism and clearance of T4 in the rat).

The incidence of observations noted at gross necropsy was low and there was no evidence of frank organ damage. On histopathologic examination of tissues, relatively mild findings occurred in both the control and treated groups. Potential test article?related histologic changes were identified in the liver and thyroid glands but these would not be considered indicative of frank toxicity. These organs were examined microscopically in all groups at both necropsies. The liver changes in male rats at the 90-day necropsy (Study Week 13) were characterized as minimal hepatocellular vacuolation and occurred in 10% of control males and ~50% of the males at 100, 300 and 1000 mg/kg/day. Minimal hepatocellular vacuolation was also detected in females in the control and test article treated groups without a clear dose response (3 to 4/10 animals per group) but, mild and moderate vacuolation was detected in females only in the 300 (1/10) and 1000 mg/kg/day (2/10) dose groups. Minimal to mild hepatocellular hypertrophy was also detected only in the 1000 mg/kg/day group (5/10) females. Minimal thyroid follicular cell hypertrophy was detected 1/10, 1/10, 5/10 and 7/10 males in the control, 100, 300 and 1000 mg/kg/day groups, respectively and in 4/10 and 3/10 females in the 300 and 1000 mg/kg/day groups respectively. In addition, mild thyroid follicular hypertrophy was detected in 4/10 females in the 1000 mg/kg/day group. The histologic changes in the liver were accompanied by an increase in liver weight. In contrast there were no statistically significant changes in thyroid weight (absolute, relative to body weight and relative to brain weight). At study week 13, mean liver weights in all dose levels of both sexes (absolute, relative to body weight and relative to brain weight) were increased compared to the male and female control means (p<0.05). The increases in liver weight were a result of a microsomal enzyme inducing effect1 and were not typically considered indicative of toxicity in absence of frank organ damage. The reversible histologic changes (vacuolation and hypertrophy) are often found to accompany increased liver weight caused by liver enzyme induction. At week 17, the liver changes (weight and histology) had at least partially, if not fully, resolved in all treated groups without delayed or long-term toxic effects. The histologic changes in the thyroid had also nearly completely resolved except in the 1000 mg/kg/day group females, where partial recovery occurred.

Increases in mean prostate weight were noted in the 1000 mg/kg/day group males at the primary necropsy. However, the increases in prostate weight were probably not of toxicological significance since the increases did not persist to the recovery period, there were no correlating histologic findings and no change in sperm production.

HBCD was detected in the adipose tissue of male and female rats treated with 1000 mg/kg/day for up to 90 days. Isomer-specific analysis showed that the relative isomer concentrations in adipose tissue at all time points were alpha>>gamma>beta which is in contrast to the test article composition (gamma>>alpha>beta). Steady state levels were achieved by study day 27. Levels in male and female rats were similar at all time points and declined during the recovery period.

All the test article-related changes at 100 and 300 mg/kg/day were mild, reversible, generally secondary to hepatic enzyme induction (which is an adaptive not a toxic change) and without effect on the clinical condition of the animals. The additional findings observed at 1000 mg/kg/day (increased gamma glutamyltransferase and additional increases in the size of the liver and prostate), were also reversible, not associated with specific target organ damage or diminished function and were, therefore, probably of limited, if any, toxicologic significance. On this basis the no-observed-adverse-effect level (NOAEL) of HBCD administered to CrI:CD®(SD)IGS BR rats by gavage in corn oil for 90 days is 1000 mg/kg/day.

Conclusions

Input optional information or further explain the contents of a particular section, much as is done in the "Discussion" portion of a publication in academic journals.

The no-observed-adverse-effect level (NOAEL) of HBCD administered to Crl:CD®(SD)IGS BR rats by gavage in corn oil for 90 days is 1000 mg/kg/day, the highest dose tested.

Data Quality

Reliability High

Denote the reliability of data, at the discretion of the person preparing the robust su

Data Reliability Remarks Add comments about how reliability of data was determined, or add re

This study was performed according to current guideline under good laboratory practices by laboratory with considerable experience in this area.

Reference

Cite the full reference for the critical study on which the robust study summary is based. List other appropriate references that support this summary and, have been reviewed for

>> Remarks

Chengelis, C. An Oral (Gavage) 90 Day Toxicity Study of HBCD in Rats. Laboratory Study No. WIL-186012. WIL Research Laboratories, Inc., Ashland, Ohio. 2001.

General

Add any statement that doesn't fit into any of the other f

Sponsored by the American Chemistry Council's Brominated Flame Retardant Industry Panel (BFRIP). Sponsor ID Albemarle Corporation Create Date 4:6:01 Study Number CAS Number Cyclododecane, 1,2.5.6,9,10-hexabromo-Consortia ID CMA Brominated Flame Retardant Industry Panel (BFRIP) Completed: **Revision Date: Test Substance** 12/5/01 The test article was a commercial HBCD product ("Hexabromid S") produced at one time by BASF in Remarks Germany. BASF no longer manufactures HBCD. **Chemical Category** Method >> Method/Guideline followed Not specified. >> GLP | Unknown >> Year study performed | 1969| Were Good Laboratory Practices followed in the st >> Species rat >> Strain Mammal strai Sprague-Dawley >> Sex | Both

| >> Number of males per dose | 10 | >> Number of females per dose | 10 |
|--|---------------------------|---|------------------------|
| >> Route of Administration | Oral | | |
| >> Exposure Period | 28 | | |
| Duration of study in days (for e | xample, 28 days, 90 d | | |
| >> Frequency of treatment | Daily | | · |
| Number of doses per day, weel | k, etc. This is particula | rly relevant for inhalation experime | ents 6hrs/day, 5 d |
| >> Doses 0, 1, 2.5, 5% of the die | et. | | |
| List all doses used in te | | | |
| >> Control Group Yes | | | |
| Concurrent contro | | | |
| >> Post observation period No | one. | | |
| Length of time animals observe | ed after last d | | |
| >> Statistical Method See Re | emarks. | | |
| Cite statistical methods use | | | |
| Remarks fo | or Method | | |
| 2.5 and 5% | of the diet for 28 days. | in Sprague-Dawley rats (10/sex/gro Doses calculated from the actual boo), 2410, and 4820 mg/kg body weight | ly weights and food |
| Results | | | |
| >> NOAEL Precision = | | | |
| >> NOAEL dose | 1000 >> Ut | iit mg/kg in feed | |
| >> NOAEL Effec | Increase in liver weight | n the absence of pathology and clinic | cal chemistry changes. |
| (e.g., decrease in body weight, organ | | | |

| > LOAEL Precision = | | | | |
|---|--------------------|-----------------|-------------------------------|------|
| > LOAEL dose | 5000 | >> Unit | mg/kg in feed | |
| > LOAEL Effect | Decrease in | boudy weight a | t a dose level of 5% in the d | iet. |
| e.g., decrease in body reight, organ | | | | |
| > Actual dose received by | y dose level by s | ex (if availa | bl | |
| 0, 940, 2410, and 4820 m | g/kg body weight/ | day | | |
| > Toxic response A brie | ef narrative desc | ribing toxic re | sponse or effects, by dos | |
| See Re | esults Remarks. | | | |
| > Statistical results Not | e statistical resu | ults, with appr | ppri ρ value | |
| | | | | |

Results Remark Provide at a minimum qualitative descriptions of elements where dose effect related observations of elements were seen:

HBCD ("Hexabromid S") was tested in Sprague-Dawley rats (10/sex/group) at doses of 0, 1, 2.5 and 5% of the diet for 28 days. Doses calculated from the body weights and food consumption are 0, 940, 2410, and 4820 mg/kg body weight/day.

No clinical signs were observed at the 1% dose levels. No significant change in mean body weight between the control and the 1 and 2.5% dose levels. The mean liver weights (absolute and relative to body weight) were different from the control mean (increased) at all dose levels, but no microscopic pathology was detected. Thyroid hyperplasia was reported in some animals at all doses, as was "very slight numerical development of the follicles and ripening follicles in the ovaries of females" at the high dose (4820 mg/kg/d). No gross or microscopic changes were detected in any other organ, and no change was detected in clinical chemistry tests.

The report concluded that "The increased liver weight must be attributed to hyperactivity; hypermetabolism as a result of increased thyroid activity appears probable in view of the observations of the thyroid". Therefore, the increased liver weights were not pathologic: there

were no microscopic lesions detected on histopathology and no change in clinical chemistry values. Recent work on the relationship of liver weight, microsomal enzyme induction, and histological change in rat toxicology studies has been published (Amacher et al, Food and Chemical Toxicology, 36, 831-839, 1998). This paper concluded "The preponderance of data collected in these 11 studies indicates that microsomal enzyme induction was not accompanied by evidence of chemically-induced liver injury. We conclude that in the rat, both hepatomegaly and microsomal enzyme induction are benign and adaptive changes in response to certain chemicals that stimulate the hepatic drug metabolizing enzyme system."

Conclusions

Input optional information or further explain the contents of a particular section, much as is done in the "Discussion" portion of a publication in academic journals.

The NOAEL in this 28-day study was 1% "Hexabromid S" in the diet. Based on body weights and food consumption data this dose is equivalent to 940 mg/kg body weight/day.

Data Quality

Reliability Reasonable

Denote the reliability of data, at the discretion of the person preparing the robust su

Data Reliability Remarks Add comments about how reliability of data was determined, or add re

This study was performed by a laboratory with considerable experience. However, the study was performed approximately 30 years ago using an HBCD product no longer manufactured as test article, and is not up to today's standards. The fact that recently conducted repeated dose studies with HBCD provided comparable results lends credence to the results of this study.

Reference

Cite the full reference for the critical study on which the robust study summary is based. List other appropriate references that support this summary and, have been reviewed for

>> Remarks

Zeller H and Kirsch P (1969) Hexabromocyclododecane: 28-day feeding trials with rats. BASF (unpublished laboratory report).

General

Add any statement that doesn't fit into any of the other f

The doses and results of this study are improperly reported by the Swedish Chemicals Inspectorate KEMI in the 1999 draft EU risk assessment of HBCD and in reports to the OECD SIDS programme. KEMI reports the doses as 0, 500, 1250 and 2500 mg/kg body weight/day (basis for conversion not given), and that the low-adverse-effect-level was 500 mg/kg (the 1% in the diet dose). Albemarie Corporation Create Date Gyclododecane 1.2.5.6.9.10-hexabromo-Study Number CMA Brominated Flame Retardant Industry Panel (BFRIP) Completed: **Revision Date:** 12/6/01 **Test Substance** The test article was a commercial HBCD product ("Hexabromid S") produced at one time by BASF in Remarks Germany. BASF no longer manufactures HBCD. **Chemical Category** >> Method/Guideline followed >> GLP Unknown >> Year study performed | 1970| Were Good Laboratory Practices followed in the st >> Strain Mammal strai Sprague-Dawley

This study was sponsored and performed by BASF.

Sponsor ID

CAS Number

Consortia ID

Method

Not known.

>> Species rat

| >> Sex Both | | |
|--|--|--|
| >> Number of males per dose 20 >> Number of females per dose 20 | | |
| >> Route of Administration Oral | | |
| >> Exposure Period 90 | | |
| Duration of study in days (for example, 28 days, 90 d | | |
| >> Frequency of treatment Daily | | |
| Number of doses per day, week, etc. This is particularly relevant for inhalation experiments 6hrs/day, 5 d | | |
| >> Doses 0, 0.16, 0.32, 0.64 and 1.28% of the diet | | |
| List all doses used in te | | |
| >> Control Group Yes | | |
| Concurrent contro | | |
| >> Post observation period 42 days. | | |
| Length of time animals observed after last d | | |
| >> Statistical Method See Remarks. | | |
| Cite statistical methods use | | |
| Remarks for Method | | |
| HBCD ("Hexabromid S") was tested in Sprague-Dawley rats at doses of 0, 0.16, 0.32, 0.64 and 1.28% of the diet for 90 days. Doses calculated on the actual body weights and food consumption in this study reveals: 0, 120, 240, 470 and 950 mg/kg body weight/day. | | |
| Results | | |
| >> NOAEL Precision = | | |
| >> NOAEL dose 1280 >> Unit mg/kg in feed | | |

| >> NOAEL Effec | Increase in liver weight in the absence of pathology or clinical chemistry changes. | | |
|---------------------------------------|---|--|--|
| (e.g., decrease in b weight, organ | ody | | |
| >> LOAEL Precisio | n > | | |
| >> LOAEL dose | 1280 >> Unit mg/kg in feed | | |
| >> LOAEL Effect | See Remarks. | | |
| (e.g., decrease in b weight, organ | ody | | |
| >> Actual dose rec | eived by dose level by sex (if availabl | | |
| 0, 120, 240, 470 a | nd 950 mg/kg body weight/day | | |
| >> Toxic response | A brief narrative describing toxic response or effects, by dos | | |
| | See Remarks. | | |
| >> Statistical resul | s Note statistical results, with appropri ρ value | | |
| See Results Rema | arks. | | |
| Results Remark | Provide at a minimum qualitative descriptions of elements where dose effect related observations of elements were seen: | | |
| | | | |

HBCD ("Hexabromid S") was tested in Sprague-Dawley rats at doses of 0, 0.16, 0.32, 0.64 and 1.28% of the diet for 90 days. Doses calculated on the actual body weights and food consumption in this study reveals: 0, 120, 240, 470 and 950 mg/kg body weight/day.

Doses up to 0.64% (470 mg/kg/d) produced no adverse clinical signs, no change in body weight, and no change in clinical chemistry results. An increase in the relative liver to body weight ratio was found, and was accompanied by fatty accumulation but no other histologically discernible changes were detected in the liver. The pathology report states that although fat ("lipid phanerosis") was visible microscopically in the liver of treated rats, this change was not accompanied by any pathology and could not be defined as "fatty liver". Further, no histological changes were found in any other organ. The report states that in the "absence of detectable clinico-chemical disturbances or histological changes of the vital organs, it was concluded that

the increased liver weight and the fat deposits, both of which were largely reversible when administration of Hexabromid S was stopped, were the result of a temporary increase in the activity of the liver."

Conclusions

Input optional information or further explain the contents of a particular section, much as is done in the "Discussion" portion of a publication in academic journals.

The highest dose tested in the BASF 90 d study, 1.28% of the diet (950 mg/kg body weight/day), is the no adverse effect level or NOAEL.

Data Quality

Reliability Reasonable

Denote the reliability of data, at the discretion of the person preparing the robust su

Data Reliability Remarks Add comments about how reliability of data was determined, or add re

This study was performed by a laboratory with considerable experience. However, the study was performed approximately 30 years ago using an HBCD product no longer manufactured as test article, and is not up to today's standards. The fact that recently conducted repeated dose studies with HBCD provided comparable results lends credence to the results of this study.

Reference

Cite the full reference for the critical study on which the robust study summary is based. List other appropriate references that support this summary and, have been reviewed for

>> Remarks

Zeller H and Kirsch P (1970) Hexabromocyclododecane: 90-day feeding trials with rats. BASF (unpublished laboratory report).

<u>General</u>

Add any statement that doesn't fit into any of the other f

This study was sponsored and performed by BASF.

The Swedish Chemicals Inspectorate (KEMI) improperly reported the results of this study and incorrectly converted the doses from a percentage of the diet to mg/kg body weight in 1999 draft EU risk assessment on HBCD and in reports to the OECD SIDS programme. KEMI converted the dietary doses to 0, 80, 160, 320 and 640 mg/kg body weight/day (basis for conversion not given). KEMI also stated that the low-adverse-effect level (LOAEL) in this

study was 80 mg/kg (the 0.16% in the diet dose level, and that a no effect level (NOEL) was not determined in any of the subchronic studies conducted to date, including the 1997 Rat 28-Day Study.

The results of the BASF 90 day study do not indicate that an adverse effect was produced at 0.16% of the diet. Further, the results indicate no adverse effect was produced at the highest dose tested, 1.28% of the diet. The BASF pathology report clearly states that although fat ("lipid phanerosis") was visible microscopically in the liver of treated rats, this change was not accompanied by any pathology and could not be defined as "fatty liver". Therefore, not even the highest dose tested in the 90-day study can be defined as the low adverse effect level (LOAEL). The highest dose tested in the BASF 90 d study, 1.28%, is more accurately defined as the no adverse effect level or NOAEL.

Toxicity End point: Toxicity in Vitro (Gene Mutations)

| era High Pi | roduction | volume (HPV) I rack | i visity ili vido (c | one mutations, |
|----------------------|--|---|--|---|
| Spensor ID | 1100021 | Albemarle Corporation | | Create Date 4/6/01 |
| CAS Number | 3194556 | Cyclododecane. 1,2,5.6,9,10-hexabron | 10- | Study Number |
| Consortia ID | 1101012 | CMA Brominated Flame Retardant Ind | ustry Panel (BFRIP) | Completed: |
| est Substance | | | | Revision Date: 12/5/01 |
| Remarks | commercial pro Chemical Corpo homogeneity. | was a composite of equal parts of the duct produced by Albemarle Corporation. The test article composite with the results of the analysis indicated mponents: HBCD beta isomer 8.5% | ration, Dead Sea Br was analyzed for ch the test article was | omine Group, and Great Lake paracterization and homogeneous and contained |
| ethod | | | | |
| >> Method/Guidelin | | | | |
| EPA OPPTS Met | thod 870.5375 In | vitro Mammalian Chromosome Abe | erration Test | |
| >> Test Type | | | | |
| Cytogenetic assa | У | | | |
| >> System of Testi | ng Non-bacteria | | | |
| >> GLP Yes | | | >> Year study pe | erformed 1996 |
| >> Species | | | | |
| Primary cultures - | human lymphod | ytes | | |
| >> Metabolic Activa | ation | | | |
| Arochlor 1254-ind | uced rat liver S-9 | ; prepared from male Sprague-Dav | vley rats | |
| >> Concentration | | | | |
| Initial: 75, 250, 75 | 0, 2500 ug/ml; D | efinitive: 10, 19, 38, 75, 150, 300, 6 | 00 ug/ml | |
| | | | | |
| >> Statistical Metho | Fisher's exa | CI (est | | |
| Remarks for Met | hod | | | |
| | The test article. | Hexabromocyclododecane (HBCD) | was tested in the ir | n vitro mammalian |

cytogenetic test using human peripheral blood lymphocytes (HPBL) in both the absence and presence of metabolic activation. The assay was performed in two phases. The first phase, the initial chromosome aberration assay, was conducted to establish the dose range for testing and to evaluate the clastogenic potential of the test article. The second phase, the independent repeat chromosome aberration assay, was performed to confirm the test system response to the test article seen in the initial assay.

Toxicity End point:
Toxicity in Vitro (Gene Mutations)

| Sponsor ID | 1103021 | Albemarle Corporation | Create Date | 1.6701 |
|--------------|---------|---|--------------|--------|
| CAS Number | 3194556 | Cyclododecane. 1.2,5,6,9,10-hexabromo- | Study Number | 1 |
| Consortia ID | 1101012 | CMA Brominated Flame Retardant Industry Panel (BFRIP) | Completed: | Y |

Dimethylsulfoxide (DMSO) was the solvent of choice based on the solubility of the test article and compatability with the target cells. The test article was soluble in DMSO at ~500 mg/ml, the highest concentration tested.

Intial. In the initial chromosome aberration assay, duplicate cultures of HPBL were exposed to 9 concentrations of the test article, and to positive, solvent and negative controls. The dividing cells were harvested at ~20 hours after initiation of treatment. The maximum dose tested was 2500 ug/ml. Dose levels greater than 2500 ug/ml were insoluble in the treatment medum and not tested. Visible precipitate was observed in treatment medium at dose levels of 750 and 2500 ug/ml and was soluble but cloudy (no visible precipitate) at dose levels 75 and 250 ug/ml. The test article was soluble in treatment medium at all other dose levels tested. In the non-activated portion of the test, HPBL cells were exposed to the test article continuously for 20 hours; in the S9-activated portion of the test, HPBL were exposed to the test article for 4 hours. Metaphase cells were collected for microscopic evaluation at 20 hours after the initiation of treatment.

Second Phase. Duplicate cultures of HPBL were exposed to at least 4 concentrations of the test article, as well as solvent, positive, and untreated controls. The dose levels selected were based on the initial assay. The dividing cells were harvested at 2 time points: 20 and 44 hours after initiation of treament. HBCD was tested in the absence and presence of an Arochlor-induced S9 metabolic activation system at dose levels of 10, 19, 38, 75, 150, 300 and 600 ug/ml. The test article was soluble but cloudy at 75 ug/ml and was workable in treatment medium at dose levels 150 ug/ml and higher. The test article was soluble in treatment medium at all other concentrations tested. In the independent repeat assay, HPBL cells were exposed to the test article continuously for 20 or 44 hours in the non-activated test system and for 4 hours in the S9-activated test system. Metaphase cells were collected for microscopic evaluation in both the non-activated and S9-activated studies at 20 and 44 hours after the initiaiton of treatment.

Evaluation of Metaphase Cells. Metaphase cells with 46 centromeres were examined under oil immersion without knowledge of treatment groups. Whenever possible, a minimum of 200 metaphase spreads (100 per duplicate treatment condition) were examined and scored for chromatid-type and chromosome-type aberrations. The mitotic index was recorded as the percentage of cells in mitosis per 500 cells counted. In the delayed harvests, the percent polyploid cells was recorded per 100 metaphase cells.

Controls. Mitomycin C was used as the positive control in the non-activated study. Cyclosphosphamide was used as the positive control in the S-9 activated study. For both positive controls one dose with sufficient scorable metaphase cells was selected for analysis. The solvent vehicle for the test article was used as the solvent control at the same concentration as that found in the test article-treated groups. Growth medium or S9 reaction mixture was used in the untreated control.

Evaluation of Results. Toxic effects of treatment were based on mitotic inhibition relative to the solvent-treated control. The number and types of aberrations, the percent aberrant cells, the percentage of numerically damaged cells and the frequency of structural aberrations per cell

Toxicity End point:
Toxicity in Vitro (Gene Mutations)

| Spansor (D | 1100021 | Albemarie Corporation | Create Date | 4/6/01 |
|--------------|---------|---|--------------|--------|
| CAS Number | 3194556 | Cyclododecane. 1,2,5,6,9,10-hexabromo- | Study Number | |
| Consortia ID | 1101012 | CMA Brominated Flame Retardant Industry Panel (BFRIP) | Completed: | Y |

was reported for each treatment group.

Results

| >> | Presult Negative |
|----|---|
| >> | Cytotoxic Concentration |
| | Non-activated: toxicity at 750 ug/ml; S9-activated: toxicity at 250 ug/ml |
| >> | Genotoxic Effects Unconfirmed |

>> Statistical results

No statistically significant differences were observed between the negative, solvent and treatment groups (p>0.05, Fisher's exact test). The positive controls performed as expected.

Results Remark

In the initial assay, dose levels of 2500 ug/ml in the non-activated study and 750 and 2500 ug/ml in the S9-activated study were not analyzed from chromosome aberrations due to complete mitotic inhibition. Toxicity (mitotic inhibition) of ~56% was observed at the highest dose level (750 ug/ml) evaluated for chromosome aberrations, in the non-activated study. In the S9-activated study, 13% toxicity was observed at the highest dose level (250 ug/ml) evaluated for chromosome aberrations. No statistically significant increases in chromosome aberrations were observed in either the non-activatged or S9-activated test systems relative to the solvent control group regardless of dose level (p>0.05, Fisher's exact test).

In the independent repeat chromosome aberration assay, toxicity, as measured by mitotic inhibition, was ~55% and 94% at the 20 and 44 hour harvest, respectively, at the highest dose levels (600 and 300 ug/ml) evaluated in the non-activated studies. In the S9-activated studies, toxicity was approximately 71% and 69% at the 20 and 44 hour harvest, respectively, at the highest dose levels (300 and 600 ug/ml) evaluated. The 600 ug/ml dose level in the non-activated 44 hour harvest and in the S9-activated 20 hour harvest was not analyzed for chromosome aberrations due to an insufficient number of scorable metaphase cells. No statistically significant increases in structural chromosome aberrations were observed in either the non-activated or S9-activated studies, regardless of dose level or harvest time (p>0.05, Fisher's exact test). No statistically significant increases in numerical chromosome aberrations were observed in either the non-activated or S9-activated studies at the 44 hour harvest time, regardless of dose level (p>0.05, Fisher's exact test).

Conclusions

| Era High | roduction | volume (FIFV) I rack | , , | |
|------------------|--------------------------------|--|------------------------|------|
| Spensor ID | 1100021 | Albemarle Corporation | Create Date | 1/6: |
| CAS Number | 3t94556 | Cyclododecane, 1,2,5,6,9,10-hexabromo- | Study Number | |
| Consortia ID | 1(01012) | CMA Brominated Flame Retardant Industry Panel (BFRIP) | Completed: Y | |
| | | ative for the induction of structural and numerical chroral blood lymphocytes. | omosome aberrations in | |
| Data Quality | Reliability H | ligh | | |
| Data Reliability | r Remarks | | | |
| | | performed using current techniques, under Good Lab considerable experience performing this type of study | | • |
| Reference | | • | | |
| >> Remarks | Lymphocytes. I | chadly, E. (1996) Chromosome Aberrations in Huma Hexabromocyclododecane. Laboratory Study Numbe Associates, Inc., Rockville, MD. | | |
| <u>General</u> | | | | |
| | Study sponsore Industry Panel. | d by the Chemical Manufacturers Association Bromin | ated Flame Retardant | |

| | Odderion | volume (FIFV) Track | |
|---------------------------|--|--|--------------------------------|
| Sponsor ID | 1100021 | Albemarle Corporation | Greate Date 4.5 |
| CAS Number | (19.356) | Cyclododecane: 1,2,5,6,9,10-hexabromo- | Study Number |
| Consortia ID | 1101012 | CMA Brominated Flame Retardant Industry Pane | ef (BFRIP) Completed: |
| | | | Revision Date: |
| est Substance | | | 12/5/01 |
| Remarks | Exact composit | on of the test article is not known. | |
| | - | | |
| | | | |
| Chemical Category | | | |
| lethod | • | | |
| >> Me thod/Guideli | ne followed | | |
| Not specified | | | |
| >> Test Type | A STATE OF THE STA | | |
| Ames test | | | |
| >> System of Testi | ng Bacterial | | |
| >> GLP Unknown | 774100 | >> Yea | ar study performed 1976 |
| | | | |
| >> Species | | | |
| Salmonella typhir | | | |
| >> Metabolic Activ | | | |
| Arochlor induced | rat liver S9 | | |
| >> Concentration | | | |
| 0, 1, 10, 50, 100, | 500, 1000, 5000 | ug/plate | |
| >> Statistical Meth | od Not known. | | |
| Remarks for Me | thod | | |
| | | almonella typhimurium (TA1535, TA1537, T | A1538, TA98 and TA100) were |
| | tested in the pre | sence and absence of a metabolic activationere 0, 1, 10, 50, 100, 500, 1000 or 5000 ug F | n system (Arochlor induced rat |

| | | n volume (Fir v) Track | |
|------------------|---|--|--|
| Sponsor (D | 110002 | Albemarle Corporation | Create Date |
| CAS Number | 319455 | Cyclododecane. 1,2,5,6,9,10-hexabromo- | Study Number |
| Consortia ID | 110101 | CMA Brominated Flame Retardant Industry Panel (BFRIP) | Completed: Y |
| ults | | | |
| Result Negati | ve | | |
| Cytotoxic Con | centration | | |
| >5000 ug HBCD |)/plate with or v | vithout metabolic activation | |
| Genotoxic Effe | unconfir | ned | |
| Statistical resu | ılts | | • |
| lot known. | | | |
| | | | |
| esults Remark | | | |
| clusions | | | |
| | tested with of performed or on Pyroguard two lots of Flactivation. In Anonymous. Sponsored bactivation testing the performance of the performance | ot mutagenic in S. typhimurium at doses up to and inclur without metabolic activation. These results are consistent this material (Ogaswara S and Hanafusa T. (1993) Red SR-103 using microorganisms; Baskin A and Phillips, M-100, Lot 53 and residue of Lot 3322 in the absence and ustrial Biotest Laboratories, Sponsored by Velsicol Ch (1979) Mutagenicity test of GLS-S6-41A. Gulf South Registry Corporation; US Environmental Protection Agency to assess the potential mutagenic effect of Compound OTS Doc #86-900000385. | tent with other Ames test's eport on mutagenicity test B. (1977) Mutagenicity of ad presence of metabolic emical Corporation; tesearch Institute, cy (1990) Ames metabolic |
| a Quality | Reliability | Acceptable | • |
| a Reliability Re | marks | | |
| | Multiple Ame | s tests performed at different test laboratories using different article have all been negative. The consistency of the confidence in the results. | |

| Sponsor ID | 1100021 | Albemarle Corporation | Create Date | 4/6/01 |
|--------------|----------|--|-------------------|--------|
| CAS Number | 319-3556 | Cyclododecane: 1,2.5.6,9,10-hexabromo- | Study Number | 2 |
| Consortia ID | 1101022 | CMA Brominated Flame Retardant Industry Panel (| BFRIP) Completed: | Y |
| Reference | | | | |
| >> Remarks | | pole, D., Newell, G., and Skinner, W. (1976) In udies for four CIBA-GEIGY Corporation compo | | 5702. |
| General | i | | | |
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| | | voidifie (i ii v) i i den | |
|---------------------|--|---|--|
| Sponsor ID | 160021 | Albemarle Corporation | Create Date 4/6/01 |
| CAS Number | 3194556 | Cyclododecane, 1,2,5,6,9,10-hexabromo- | Study Number |
| Consortia ID | 1101012 | CMA Brominated Flame Retardant Industry Panel (BFRIF | Completed: Y |
| | | | |
| | 7 | | Revision Date: |
| est Substance | | | 12/6/01 |
| Remarks | HBCD, obtained | from Aldrich Chemicals (Stockholm, Sweden) | |
| hemical Category | 7 | | |
| ethod | annad | | |
| | II <i>(</i> - | • | |
| >> Method/Guide | ······································ | | |
| Non standard to | est methodology | | |
| >> Test Type | | | |
| Mammalian cells | s in culture (Sp5 ar | nd SPD8 duplication cell lines) | |
| >> System of Tes | ting Non-bacterial | | |
| >> GLP No | | >> Year stud | y performed 1999 |
| >> Species | | | |
| Not known. | | | |
| >> Metabolic Acti | vation | | |
| None | | | |
| >> Concentration | The second secon | | |
| See results. | CHARLES HARRISTON OF THE STATE | | |
| >> Statistical Meti | hod Student's t to | est | |
| Remarks for Me | ethod | | |
| | five doses betwee spontaneous par protein. The mu recombination; a | ed in vitro in hamster cells (Sp5/V79 and SPD8) in a sen 2 and 20 ug/ml plus a control. The Sp5 and SP rtial duplication of the HPRT gene, resulting in a nor tants revert spontaneously to a functional HPRT ge in increase in reversion frequency is considered a p | D8 clones exhibit a n-functional HGPRT ene phenotype by positive response. |

| Spensor ID | (1009. | Albemarle Corporation | Create Date 1/6/01 |
|--|---|---|----------------------------|
| CAS Number | 3 [Q. E.) | 6 Cyclododecane, 1,2,5,6,9,10-hexabromo- | Study Number 3 |
| Consortia (D | 11019: | CMA Brominated Flame Retardant Industry Panel (BFRIP |) Completed: Y |
| | This reliabilithe system, | d as statistically significant. by of this genetic test is unknown. The reproducibility of dose-effect response, and whether a maximal two-fold bonse are also unknown. | |
| >> Result Ambigu >> Cytotoxic Cond Not known. | | | |
| >> Genotoxic Effe | cts Equivoca | | |
| >> Statistical resu See Remarks See Results Remark | cttion. | | |
| | was reported This reliabilithe system, | ith HBCD resulted in a ~ maximal 2-fold increase in revel as statistically significant. y of this genetic test is unknown. The reproducibility of dose-effect response, and whether a maximal two-fold is onse are also unknown. | the results, validation of |
| onclusions | | | |
| | | ith HBCD resulted in a ~ maximal 2-fold increase in revol I as statistically significant. The reliability of this test is | |
| Data Quality | Reliability | Unknown. | |
| Data Reliability Rer | marks | | |

| <u> </u> | | | | |
|----------------|--------------------|--|--------------|--|
| Spansor ID | 1100021 | Albemarle Corporation | Create Date | 4/6,01 |
| CAS Number | 319-1556 | Cyclododecane. 1,2.5,6,9.10-hexabromo- | Study Number | 3] |
| Consortia ID | 1101012 | CMA Brominated Flame Retardant Industry Panel (BFRIP) | Completed: | Υ |
| | results, validatio | nd predictive ability of this genetic test is unknown. The of the system, dose-effect response, and whether a ence of a positive response are unknown. | | the |
| Reference | | | | |
| >> Remarks | Helleday et al, N | flutat Res, 1999, 439(2): 137-147. | | en e |
| <u>General</u> | | | | |
| | | | | |
| | | | | |

Toxicity End Point:

| EPA High Pr | oduction | Volume (HP | V) Track | Toxicity in Vivo (Ch | romosomal Aber | rations) |
|---------------------------|--|---|--|---|--|--|
| Spansor ID | 1200021 | Albemarle Corporati | on | | Create Date | 1/6/01 |
| CAS Number | 3 ; 0 4 5 % (5 | Cyclododecane: 1,2, | 5,6,9,10-hexabromo- | | Study Number | 1 |
| Consortia (D | 1101012 | CMA Brominated Fla | ime Retardant Indus | try Panel (BFRIP) | Completed: | Υ |
| Test Substance Remarks | commercial prod Lakes Chemica homogeneity. T | was a composite of duct produced by Al Corporation. The t | bemarle Corporat test article compos alysis indicated th | ion, Dead Sea Bro site was analyzed e test article was l | promocyclodod omine Group, a for characteriza homogeneous | nd Great ation and and contained |
| | isomer 79.1%. | mponents: HBCD be | ata isomer 6.5%, i | TBCD alpha isome | er 6.0%, HBCD | gamma |
| Chemical Category | | | | | | |
| <u>Method</u> | | | | | | |
| >> Method/Guidelin | ne followed | | | | | |
| OECD Method 47 | 4 | | | | | |
| >> Test Type | · | | • | | | |
| Micronucleus ass | ay | | | | | |
| >> GLP Yes | | | >> | Year study perfe | ormed 2000 | |
| >> Species | | | | | | |
| mouse | | | | | | |
| >> Strain Mammal | strai NMRI | | Welfur and the state of the sta | Andrew Colonia de La companya de la Colonia | | |
| >> Sex M | | | | | | 1 |
| >> Number of male | s per dose | 5 | >> Number of f | emales per dose | | 0 |
| >> Route of Admini | stration | | | | The state of the s | ······································ |
| Intrperioneal | | | | | | |
| >> Doses 0, 500, 1 | 000, 2000 mg/kg | | | | | |
| >> Exposure period | Two doses a | dministered 24 hrs a | apart. | | | |
| >> Statistical Metho | Wilcoxon te | st | | | | |

Remarks for Method

Toxicity End Point:
Toxicity in Vivo (Chromosomal Aberrations)

| Sponsor ID | 1100021 | Albemarle Corporation | Create Date | 1/6/01 |
|--------------|---------|---|--------------|--------|
| CAS Number | 3194556 | Cyclododecane, 1,2,5,6.9,10-hexabromo- | Study Number | 1 |
| Consortia ID | 1101012 | CMA Brominated Flame Retardant Industry Panel (BFRIP) | Completed: | Y |

Hexabromocyclododecane (HBCD) was tested for clastogenicity and for the ability to induce spindle poison effects in NMRI mice (Charles River Deutschland GmbH) using the micronucleus method. HBCD, dissolved in DMSO, was administered twice intraperitoneally with a 24-hr interval between doses to male mice (n=5/group) at dose levels of 500, 1000 or 2000 mg/kg body weight in a volume of 4 ml/kg. DMSO (the vehicle) was administered to male mice by the same route and frequency. Cyclophosphamide was used as a positive control for clastogenic effects. Vincristine was used as a positive control for induction of spindle poison effects. Animals in the positive control groups were treated only once.

The animals were sacrificed and the bone marrow of the two femora prepared 24 hours after the second administration. After staining, 2000 polychromatic erythrocytes were evaluated per animal and investigated for micronuclei. The normocytes with and without micronuclei occurring per 2000 polychromatic erythrocytes were also counted.

Results

>> Effects on Mitosi

PCE/NCE 0, 500, 1000, 2000 mg/kg = 3.74, 2.89, 2.67, 2.49, respectively.

>> Genotoxic Effects Negative

>> Statistical results

No statistical differences between the treatment and vehicle control group were observed (p<=0.05).

Results Remark

The two intraperitoneal administrations of DMSO in a volume of 4 ml/kg body weight led to 1.4% polychromatic erythrocytes containing micronulei. In the 2000 mg HBCD/kg body weight group, 0.9% micronuclei were found. In the 1000 and 500 mg HBCD/kg body weight groups, 1.0 and 1.1% micronuclei were detected. The two positive control substances performed as expected.

The number of normochromatic erythrocytes containing micronuclei did not diffeer to any appreciable extent in the negative control or various dose groups.

Conclusions

HBCD treatment did not increase numbers of micronuclei. The number of normochromatic or polychromatic erythrocytes containing small micronuclei did not deviate from the vehicle control value and was within the historical control range. Large micronuclei were not observed. HBCD had no chromosome-damaging (clastogenic) effect in this study and did not impair chromosome distribution during mitosis.

EPA High Production Volume (HPV) Track Toxicity End Point: Toxicity In Vive (Chromosomal Aberrations)

| C. 7. 1g | Treatment votation (111 v) Tract Toxicity in vivo (circumstantial Abeliations) | | | | | |
|--------------------|--|---|--|--------|--|--|
| Sponsor ID | 1:00021 | Albemarle Corporation | Create Date | 47670) | | |
| CAS Number | 3194556 | Cyclododecane: 1,2,5.6,9,10-hexabromo- | Study Number | | | |
| Consortia ID | 1101012 | CMA Brominated Flame Retardant Industry Panel (BFRIP) | Completed: | Υ | | |
| Data Quality | Reliability Hi | gh | | | | |
| Data Reliability R | emarks | | | | | |
| | This study was performed according to current guidelines under Good Laboratory Practices by an experienced laboratory. | | | | | |
| Reference | | | | | | |
| >> Remarks | Engelhardt, G and Hoffmann, H. (2000) Cytogenetic Study in vivo with Hexabromocyclododecane in the Mouse Micronucleus Test After Two Intraperitoneal Adminstrations. Laboratory Project Identification: 26M0100/004018. Experimental Toxicology and Ecology, BASF Aktiengesellschaft, Ludwigshafen, Germany. | | | | | |
| <u>General</u> | i. | | | | | |
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